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(54) Title: METHOD FOR PRODUCING CAROTENOID COMPOUNDS IN PLANTS

(57) Abstract: Methods are provided for producing plants and seeds having altered carotenoid compositions by transforming host plants with constructs having a transcriptional initiation region from a gene expressed in a plant seed, a plastid transit peptide, a DNA sequence derived from at least one carotenoid biosynthesis gene coding region, and a transcriptional termination region. The methods find particular use in increasing the carotenoid content in oilseed plants.

A. CLASSIFICATION OF SUBJECT MATTER
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Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

C12N15/53

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Category •	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO 98 06862 A (CALGENE INC (US)) 19 February 1998 (1998-02-19) the whole document, especially page 14, lines 1-6 and page 18, lines 16-22	1-16,19, 20
X	WO 99 07867 A (CALGENE LLC) 18 February 1999 (1999-02-18) the whole document	1-16,19, 20
X	WO 96 13149 A (AMOCO CORP) 9 May 1996 (1996-05-09) cited in the application page 7, line 30 -page 10, line 19 page 23, line 15 -page 36, line 21	1-4, 7-10,14, 19,20
(WO 99 55889 A (DU PONT) 4 November 1999 (1999-11-04) the whole document	17,19-21

	<u>X</u>	Further docum	nents are listed in the	continuation of box C.
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Patent family members are listed in annex.

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8 May 2002 21/05/2002

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INTERNATIONAL SEARCH REPORT

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Category °	Citation of document, with indication, where appropriate, of the relevant passages			
X .	BUCKNER BRENT ET AL: "The y1 gene of maize codes for phytoene synthase." GENETICS, vol. 143, no. 1, 1996, pages 479-488, XP001040499 ISSN: 0016-6731 abstract		17,18,22	
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INTERNATIONAL SEARCH REPORT

Information on patent family members

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(57) Abstract: Methods are provided for producing plants and seeds having altered carotenoid compositions by transforming host plants with constructs having a transcriptional initiation region from a gene expressed in a plant seed, a plastid transit peptide, a DNA sequence derived from at least one carotenoid biosynthesis gene coding region, and a transcriptional termination region. The methods find particular use in increasing the carotenoid content in oilseed plants.

METHODS FOR PRODUCING CAROTENOID COMPOUNDS, AND SPECIALITY OILS IN PLANT SEEDS

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This application is a continuation-in-part of Application Serial No. 09/023,587, filed February 13, 1998, and a continuation-in-part of Application Serial No. 09/130,549, filed August 6, 1998, which is a continuation-in-part of Application Serial No.08/908,758 filed August 8, 1997 which claims the benefit of the filing date of provisional Application Serial No.60/024,145 filed August 9, 1996.

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FIELD OF THE INVENTION

The invention relates to genetic modification of plants, plant cells and seeds, particularly altering carotenoid biosynthesis, and fatty acid composition.

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BACKGROUND OF THE INVENTION

Carotenoids are pigments with a variety of applications. They are yellow-orange-red lipids which are present in green plants, some molds, yeast and bacteria. Carotenoid hydrocarbons are referred to as carotenes, whereas oxygenated derivatives are referred to as xanthophylls. The carotenoids are part of the larger isoprenoid biosynthesis pathway which, in addition to carotenoids, produces such compounds as chlorophyll and tocopherols, Vitamin E active agents. The carotenoid pathway in plants produces carotenes, such as α - and β -carotene, and lycopene, and xanthophylls, such as lutein.

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The biosynthesis of carotenoids involves the condensation of two molecules of the C_{20} precursor geranyl PP_i to yield the first C_{40} hydrocarbon phytoene. In a series of sequential desaturations, phytoene yields lycopene. Lycopene is the precursor of the

cyclic carotenes, β -carotene and α -carotene. The xanthophylls, zeaxanthin and lutein are formed by hydroxylation of β -carotene and α -carotene, respectively.

β-carotene, a carotene whose color is in the spectrum ranging from yellow to orange, is present in a large amount in the roots of carrots and in green leaves of plants. β-carotene is useful as a coloring material and also as a precursor of vitamin A in mammals. Current methods for commercial production of β-carotene include isolation from carrots, chemical synthesis, and microbial production.

A number of crop plants and a single oilseed crop are known to have substantial levels of carotenoids, and consumption of such natural sources of carotenoids have been indicated as providing various beneficial health effects. The below table provides levels of carotenoids that have been reported for various plant species.

CAROTENOID CONTENTS OF VARIOUS CROPS (μg/g)

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Crop	Beta-Carotene	Alpha-Carotene	Lycopene	Lutein	Total
Carrots	30-110	10-40	0-0.5	0-2	65-120
Pepper (gr)	2	-	-	2	8
Pepper (red)	15	1	-	-	200
Pumpkin	16	0.3	tr	26	100
Tomato	3-6	-	85	-	98
Watemelon	1	tr	19	-	25
Marigold peta	ls 5	4	-	1350	1500
Red palm oil	256	201	8	-	545

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The pathway for biosynthesis of the carotenoids has been studied in a variety of organisms and the biosynthetic pathway has been elucidated in organisms ranging from bacteria to higher plants. See, for example, Britton, G. (1988) Biosynthesis of carotenoids, p. 133-182, In T.W. Goodwin (ed.), Plant pigments, 1988. Academic

Press, Inc. (London), Ltd., London. Carotenoid biosynthesis genes have also been cloned from a variety of organisms including Erwinia uredovora (Misawa et al. (1990) J. Bacteriol. 172:6704-6712; Erwinia herbicola (Application WO 91/13078, Armstrong et al. (1990) Proc. Natl. Acad. Sci., USA 87:9975-9979); R. capsulatus (Armstrong et al. (1989) Mol. Gen. Genet. 216:254-268, Romer et al. (1993) Biochem. Biophys. Res. Commun. 196:1414-1421); Thermus thermophilus (Hoshino et al. (1993) Appl. Environ. Microbiol. 59:3150-3153); the cyanobacterium Synechococcus sp. (Genbank accession number X63873). See also, application WO 96/13149 and the references cited therein.

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While the genes have been elucidated, little is known about the use of the genes in plants. Investigations have shown that over expression or inhibition of expression of the plant phytoene synthase (Psy1) gene in transgenic plants can alter carotenoid levels in fruits. See, Bird et al. (1991) Biotechnology 9:635-639; Bramley et al. (1992) Plant J. 2:343-349; and Fray and Grierson (1993) Plant Mol. Biol. 22:589-602. Further, as reported by Fray et al. (1995) The Plant Journal 8:693-701, constitutive expression of a fruit phytoene synthase gene in transgenic tomatoes causes dwarfism by redirecting metabolites from the gibberellin pathway.

Application WO 96/13149 reports on enhancing carotenoid accumulation in storage organs such as tubers and roots of genetically engineered plants. The application is directed towards enhancing colored native carotenoid production in specific, predetermined non-photosynthetic storage organs. The examples of the application are drawn to increasing colored carotenoids in transformed carrot roots and in orange flesh potato tubers. Both of these tissues are vegetative tissues, not seeds, and natively have a high level of carotenoids.

Carotenoids are useful in a variety of applications. Generally, carotenoids are useful as supplements, particularly vitamin supplements, as vegetable oil based food products and food ingredients, as feed additives in animal feeds and as colorants.

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Specifically, phytoene finds use in treating skin disorders. See, for example, U.S. Patent No. 4,642,318. Lycopene, α - and β -carotene are used as food coloring agents. Consumption of β -carotene and lycopene has also been implicated as having preventative effects against certain kinds of cancers. In addition, lutein consumption has been associated with prevention of macular degeneration of the eye.

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Plant oils are useful in a variety of industrial and edible applications. Novel vegetable oils compositions and/or improved means to obtain oils compositions, from biosynthetic or natural plant sources are needed. Depending upon the intended oil use, various different fatty acid compositions are desired. The demand for modified oils with specific fatty acid compositions is great, particularly for oils high in oleic acid. See, Haumann, B. F. (1996) *INFORM* 7:320-334. As reported by Haumann, the ideal frying oil would be a low-saturate, high oleic and low linolenic oil. Furthermore, studies in recent years have established the value of monounsaturated fatty acids as a dietary constituent.

Attempts have been made over the years to improve the fatty acid profiles of particular oils. For example, the oxidative stability of vegetable oil is related to the number of double bonds in its fatty acids. That is, molecules with several double bonds are recognized to be more unstable. Thus, scientists have attempted to reduce the content of α -linolenic acid in order to improve shelf life and oxidative stability, particularly under heat.

It is apparent that there is needed a method for producing significant levels of carotenoid compounds in crop plants and particularly in plant seeds. It would additionally be beneficial to alter the fatty acid content of the plants and seeds. Such altered seed products would be useful nutritionally as well as provide a source for producing more stable oils. There is no report of methods to substantially altering the levels and composition of carotenoids produced in a plant seed, particularly with respect

to increasing the level of production of carotenoids. There is therefore needed, a useful method for altering carotenoid levels in plants, particularly seeds, and for producing oils with modified carotenoid composition and/or content.

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SUMMARY OF THE INVENTION

Transformed plants, plant cells and seeds having altered carotenoid levels and/or modified fatty acid compositions are provided. The plants, plant cells and seeds are transformed with at least one carotenoid biosynthesis gene, or a combination thereof. Methods for making and using the transformed compositions of the invention are also provided. Methods find use in altering carotenoid levels in plants, particularly seeds, as well as increasing particular compounds for molecular farming, such as for production of particular carotenoids. At the same time, the transformed compositions, particularly seeds, provide a source of modified oils, which oils may be extracted from the seeds in order to provide an oil product comprising a natural source of various carotenoids, carotenoid mixtures. In a particular aspect of the present invention, transformed seed can provide a source for particular carotenoid compounds and/or for modified specialty oils having altered carotenoid compositions and/or altered fatty acid composition, particularly having increased levels of oleic acid and decreased levels of linoleic and linolenic acids.

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BRIEF DESCRIPTION OF THE FIGURES

Figure 1 shows the nucleotide sequence of the SSU/crtB fusion sequence, SEQ ID No. 1.

Figure 2 presents constructs for expression of carotenoid biosynthesis genes in plant seeds. Figure 2A shows plasmid pCGN3390 which con

tains the napin promoter operably linked to the SSU/crtB sequence. Figure 2B shows plasmid pCGN3392which contains the napin promoter operably linked to the SSU/crtE sequence. Figure 2C shows plasmid pCGN9010 which contains the napin promoter operably linked to the SSU/crtI sequence. Figure 2D shows plasmid pCGN9009 which contains the napin promoter operably linked to the SSU/crtB sequence and the napin promoter operably linked to the SSU/crtI sequence. Figure 2E shows plasmid pCGN9002 which contains the napin promoter operably linked to the SSU/crtB sequence and the napin promoter operably linked to an antisense epsilon cyclase sequence. Figure 2F shows plasmid pCGN9017 which contains the napin promoter operably linked to the SSU/crtB sequence and the napin promoter operably linked to an antisense beta cyclase sequence. Figure 2G shows plasmid pCGN6204 which contains the napin promoter operably linked to the SSU/crtB sequence and the napin promoter operably linked to the SSU/crtW sequence. Figure 2H shows plasmid pCGN6205 which contains the napin promoter operably linked to the SSU/crtB sequence and the napin promoter operably linked to the crtZ sequence. Figure 2I shows plasmid pCGN6206 which contains the napin promoter operably linked to the SSU/crtB sequence, the napin promoter operably linked to the crtW sequence and the napin promoter operably linked to the crtZ sequence. Figure 2J provides a schematic diagram of the corn expression construct pCGN9039.

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Figure 3 shows the results of analyses of saponified samples for control seeds.

Figure 4 shows the results of analyses of saponified samples for pCGN3390 transformed seeds.

Figure 5 shows a graph of the fatty acid analysis in pCGN3390 transformed seeds and demonstrates that the increase in 18:1 fatty acids correlates with a decrease in 18:2 and 18:3.

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Figure 6 shows a graph of the fatty acid analysis in pCGN3390 transformed seeds and demonstrates that the increase in 18:1 correlates with an increase in both 18:0 and 20:0, but little effect is seen in 16:0.

Figure 7 shows a graph of the fatty acid analysis in pCGN3390 transformed seeds and demonstrates the increase in 18:0 correlates well with an increase in 20:0.

Figure 8 shows a carotenoid biosynthesis pathway.

Figure 9 provides sequence of B. napus epsilon cyclase cDNA clone 9-4, SEQ ID No. 2.

Figure 10 provides sequence of B. napus epsilon cyclase cDNA clone 7-6, SEQ ID No. 3.

Figure 11 provides sequence of a B. napus beta cyclase cDNA clone, SEQ ID No. 4.

Figure 12 provides T2 seed analysis of 3390 transformed Brassica napus plants.

Figure 13 provides T3 seed analysis of 3390 transformed Brassica napus plants.

Figure 14 provides T2 seed analysis of 9002 transformed Brassica napus plants.

Figure 15 shows the nucleotide sequence of the SSU/crtZ fusion sequence, SEQ ID No. 5, and the deduced amino acid sequence SEQ ID No. 6.

Figure 16 shows the nucleotide sequence of the SSU/crtW fusion sequence, SEQ ID No. 7, and the deduced amino acid sequence SEQ ID No. 8.

Figure 17 shows the HPLC trace for detection of xanthophylls from extractions from seed of 6204 transgenic lines.

Figure 18 provides the results of the expression of the maize phytoene synthase in *Arabidopsis* comparing the levels of B-carotene to total carotenoid levels in 9061 lines.

Figure 19 provides the complete nucleic acid sequence of the maize phytoene synthase sequence SEQ ID No. 9, and the deduced amino acid sequence, SEQ ID No. 10.

DETAILED DESCRIPTION OF THE INVENTION

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In accordance with the subject invention, methods for increasing production of carotenoid compounds, as well as for altering fatty acid compositions in a plant, particularly in plant seeds, are provided. The method involves transforming a plant cell with at least one carotenoid biosynthesis biosynthesis gene. This has the effect of altering carotenoid biosynthesis, particularly increasing the production of downstream products, as well as providing novel seed oils having desirable fatty acid compositions. A second gene can then be utilized to shunt the metabolic activity to the production of particular carotenoid, or to further alter the fatty acid composition.

Surprisingly, it has been found that transformation of a plant with an early carotenoid biosynthesis gene leads to a significant increase in the flux through the carotenoid pathway resulting in an increase in particular carotenoids. That is, there is an increase in the metabolic activity that can be further manipulated for the production of specific carotenoids. In addition, the transformed seeds may demonstrate altered fatty acid compositions as the result of the carotenoid gene expression, such as seen with the seeds described herein from plants transformed with a phytoene synthase gene.

Thus, using the methods of the invention, seeds are provided which produce high levels of particular carotenoids and/or produce specialty oils having a desired fatty acid composition. In oilseed Brassica, for example, transformation with an early carotenoid biosynthesis gene leads to seeds having significant increases in the levels of α -carotene, β -carotene and lutein. In addition, the Brassica seeds demonstrate an altered fatty acid composition and yield a vegetable oil which has increased oleic acid content and

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decreased linoleic and linolenic acid content. Thus, the transformed seed can provide a source of carotenoid products as well as modified seed oil. In this manner, modified specialty oils can be produced and new sources of carotenoids for extraction and purification are provided.

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The oils of the present invention also provide a substantial improvement with respect to stability as compared to two other major plant sources of carotenoids, marigold petals and red palm oil (mesocarp). Although instability is observed in seeds stored in air at room temperature as demonstrated by loss of approximately 20-30% of total carotenoids after 4 weeks of storage, the loss after 1-2 weeks is only 10%. Palm mesocarp, by contrast, must be processed within a day or two of harvest in order to avoid major losses of carotenoids. Furthermore, the carotenoid decomposition in the seeds of the present invention may be reduced significantly by storage of the seeds under nitrogen.

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For the production of a seed having an increase in carotenoid biosynthesis, transformation of the plant with an early carotenoid biosynthesis gene is sufficient. By early carotenoid biosynthesis gene is intended geranylgeranyl pyrophosphate synthase, phytoene synthase, phytoene desaturase, and isopentenyl diphosphate (IPP) isomerase. A variety of sources are available for the early carotenoid biosynthesis genes and for the most part, a gene from any source can be utilized. However, it is recognized that because of co-suppression, the use of a plant gene native to the target host plant may not be desirable where increased expression of a particular enzyme is desired.

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A number of early carotenoid biosynthesis genes, also referred herein as DNA sequences derived from carotenoid biosynthesis gene coding regions, have been isolated and are available for use in the methods of the present invention. See, for example:

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IPP isomerase has been isolated from: R. Capsulatus (Hahn et al. (1996) J. Bacteriol. 178:619-624 and the references cited therein), GenBank Accession Nos.

U48963 and X82627, Clarkia xantiana GenBank Accession No. U48962, Arabidopsis thaliana GenBank Accession No. U48961, Schizosaccharmoyces pombe GenBank Accession No. U21154, human GenBank Accession No. X17025, Kluyveromyces lactis GenBank Accession No. X14230;

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geranylgeranyl pyrophosphate synthase from E. Uredovora Misawa et al. (1990)

J. Bacteriol. 172:6704-6712 and Application WO 91/13078; and from plant sources,
including white lupin (Aitken et al. (1995) Plant Phys. 108:837-838), bell pepper
(Badillo et al. (1995) Plant Mol. Biol. 27:425-428) and Arabidopsis (Scolnik and
Bartely (1994) Plant Physiol. 104:1469-1470; Zhu et al. (1997) Plant Cell Physiol.
38:357-361).

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phytoene synthase from a number of sources including E. Uredovora, Rhodobacter capsulatus, and plants Misawa et al. (1990) J. Bacteriol. 172:6704-6712, GenBank Accession No. D90087, Application WO 91/13078, Armstrong et al. (1989) Mol. Gen. Genet. 216:254-268, Armstrong, G. A. "Genetic Analysis and regulation of carotenoid biosynthesis. In R. C. Blankenship, M. T. Madigan, and C. E. Bauer (ed.), Anoxygenic photosynthetic bacteria; advances in photosynthesis. Kluwer Academic Publishers, Dordrecht, The Netherlands, Armstrong et al. (1990) Proc. Natl. Acad. Sci USA 87:9975-9979, Armstrong et al. (1993) Methods Enzymol. 214:297-311, Bartley and Scolnik (1993) J. Biol. Chem. 268:27518-27521, Bartley et al. (1992) J. Biol. Chem. 267:5036-5039, Bramley et al. (1992) Plant J. 2:291-343, Ray et al. (1992) Plant Mol. Biol. 19:401-404, Ray et al. (1987) Nucleic Acids Res. 15:10587, Romer et al. (1994) Biochem. Biophys. Res. Commun. 196:1414-1421, Karvouni et al. (1995) Plant Molecular Biology 27:1153-1162, GenBank Accession Nos. U32636, Z37543, L37405, X95596, D58420, U32636, Z37543, X78814, X82458, S71770, L27652, L23424, X68017, L25812, M87280, M38424, X69172, X63873, and X60441, Armstrong, G. A. (1994) J. Bacteriol. 176:4795-4802 and the references cited therein; and,

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phytoene desaturase from bacterial sources including E. uredovora Misawa et al. (1990) J. Bacteriol. 172:6704-6712, and Application WO 91/13078 (GenBank Accession Nos. L37405, X95596, D58420, X82458, S71770, and M87280); and from plant sources, including maize (Li et al. (1996) Plant Mol. Biol. 30:269-279), tomato (Pecker et al. (1992) Proc. Nat. Acad. Sci. 89:4962-4966 and Aracri et al. (1994) Plant Physiol. 106:789), and Capisum annuum (bell beppers) (Hugueney et al. (1992) J. Biochem. 209: 399-407), GenBank Accession Nos. U37285, X59948, X78271, and X68058).

See, generally, Misawa et al. (1990) J. of Bacteriology 172:6704-6712, E.P. 0393690 B1, U.S. Patent No. 5,429,939, Bartley et al. (1992) J. Biol. Chem. 267:5036-5039, Bird et al. (1991) Biotechnology 9:635-639, and US Patent No. 5,304,478, which disclosures are herein incorporated by reference.

Transformation with an early carotenoid gene, (herein referred to as the primary gene), increases the biosynthetic activity of the carotenoid pathway, and can lead to increased production of particular carotenoids such as for example, α - and β -carotene. As described in more detail in the following examples, by expression of phytoene synthase as the primary gene, large increases in the carotenoid content generally, and particularly in the levels of α - and β -carotene, are obtained in seeds of transformed plants. Oil comprising the carotenoids so produced may be extracted from the seeds to provide a valuable source of α - and β -carotenes. Such an oil may find use as a food colorant, for example to add color to margarines, or as a food oil. An edible food oil with high α - and β -carotene levels is of interest for prevention of Vitamin A deficiency which can result in night blindness. Thus, production of transformed plants and extraction of the high α - and β -carotene oil to provide a useful food oil is particularly desirable in regions where night blindness is a widespread problem, such as in India and Asia.

In addition to high α - and β -carotene levels, levels of other carotenoids are also increased in the oils exemplified herein. For example, lutein levels are increased in seeds from plants transformed with a phytoene synthase gene, as well as in seeds from plants transformed with a GGPP synthase gene, crtE (3392), or with phytoene desaturase, crtI (9010).

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Furthermore, additional primary genes may be expressed to provide for even greater flux through the carotenoid pathway. For example, in oilseed *Brassica* seeds transformed with a phytoene synthase gene as described herein, increased levels of phytoene are observed. Thus, increasing the expression of phytoene desaturase as well as phytoene synthase may result in further increases in the levels of carotenoids, such as α- and β-carotene and lutein, produced. Such further modification of carotenoid composition is demonstrated here in transgenic plant seeds transformed with pCGN9009 for the expression of *crt*B and *crt*I genes. Additionally, plants expressing both phytoene synthase and GGPP synthase genes are desirable. Such plants may demonstrate even greater flux through the carotenoid pathway as indicated by the increased production of chlorophyll observed in plants of the present invention which have been transformed to express a GGPP synthase gene (*crt*E) in the absence of *crt*B overexpression.

Interestingly, plants expressing a GGPP synthase gene did not have significant modifications of the tocopherol content. Since GGPP is a branch point of the carotenoid, chlorophyll and tocopherol pathways in plants, these observations suggest that the next enzymatic step in tocopherol biosynthesis, catalyzed by GGPP hydrogenase, is a rate limiting step for tocopherol production. Thus, providing for increased expression of GGPP hydrogenase, alone or in conjunction with increased expression of GGPP synthase would be expected to result in an increase of flux to the tocopherol pathway.

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Also of interest are plants which are transformed to express three early carotenoid biosynthesis gens, crtB, crtE, and crtI. Plants expressing two or three different carotenoid biosynthsis genes may be produced by either transforming a plant with a construct providing for expression of the desired genes, using a multiple gene construct or by cotransformation with multiple constructs, or by crossing plants which contain the different desired genes.

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In addition to the production of the carotenoids described herein, once the biosynthetic activity has been increased by expression of the primary carotenoid biosynthesis gene or genes, the pathway can be diverted for the production of specific compounds. The diversion involves the action of at least one second gene of interest, (the secondary gene). The secondary gene can encode an enzyme to force the production of a particular compound or alternatively can encode a gene to stop the pathway for the accumulation of a particular compound. For forcing the production of a particular compound, expression of a carotenoid biosynthesis gene in the pathway for the desired carotenoid compound is used. Genes native or foreign to the target plant host may find use in such methods, including, for example, carotenoid biosynthesis genes from sources other than higher plant, such as bacteria, including Erwinia and Rhodobacter species. For stopping the pathway in order to accumulate a particular carotenoid compound, the secondary gene will provide for inhibition of transcription of a gene native to the target host plant, wherein the enzyme encoded by the inhibited gene is capable of modifying the desired carotenoid compound. Inhibition may be achieved by transcription of the native gene to be inhibited in either the sense (cosuppression) or antisense orientation of the gene.

For example, for alteration of the carotenoid composition towards the accumulation of higher levels of \(\mathcal{B}\)-carotene derived carotenoids, such as zeaxanthin, zeaxanthin diglucoside, canthaxanthin, and astaxanthin, inhibition of lycopene epsilon

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cyclase is desired to prevent accumulation of alpha carotene and its derivative carotenoids, such as lutein. In addition, overexpression of lycopene β -cyclase may be used to increase the accumulation of β -carotene derived carotenoids. Thus, antisense lycopene epsilon cyclase and lycopene β -cyclase are examples of sequences which find use in secondary gene constructs of interest in the present invention. Furthermore, in conjunction with the inhibition of lycopene epsilon cyclase, increased expression of additional secondary genes may be desired for increased accumulation of a particular beta-carotene derived carotenoid. For example, increased β -carotene hydroxylase expression is useful for production of zeaxanthin, wherease increased β -carotene hydroxylase and keto-introducing enzyme expression is useful for production of astaxanthin. Alternatively, for accumulation of lycopene, inhibition of lycopene beta cyclase or of lycopene epsilon cyclase and lycopene beta cyclase is desired to reduce conversion of lycopene to alpha- and beta-carotene.

Thus, the carotenoid pathway can be manipulated by expression of carotenoid biosynthesis genes to increase production of particular carotenoids, or by decreasing levels of a particular carotenoid by transformation with antisense DNA sequences which prevent the conversion of a selected precursor compound into the next carotenoid in the pathway.

Secondary genes of interest in the present application include but are not limited to:

β-carotene hydroxylase or crtZ (Hundle et al. (1993) FEBS Lett. 315:329-334, GenBank Accession No. M87280) for the production of zeaxanthin;

genes encoding keto-introducing enzymes, such as crtW (Misawa et al. (1995) J. Bacteriol. 177:6575-6584, WO 95/18220, WO 96/06172) or B-C-4-oxygenase (crtO; Harker and Hirschberg (1997) FEBS Lett. 404:129-134) for the production of canthaxanthin;

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crtZ and crtW or crtO for the production of astaxanthin; ε -cyclase and ε -hydroxylase for the production of lutein; ε -hydroxylase and crtZ for the production of lutein and zeaxanthin; lycopene β -cyclase (crtY) (Hugueney et al. (1995) Plant J.

8:417-424, Cunningham FX Jr (1996) Plant Cell 8:1613-1626, Scolnik and Bartley (1995) Plant Physiol. 108:1343, GenBank Accession Nos. X86452, L40176, X81787, U50739 and X74599) for increased production of β-carotene.

antisense lycopene ϵ -cyclase (GenBank Accession No. U50738) for increased production of β -carotene;

antisense lycopene ϵ -cyclase and lycopene β -cyclase for the production of lycopene;

antisense plant phytoene desaturase for the production of phytoene; etc.

In this manner, the pathway can be modified for the high production of any particular carotenoid compound of interest, or for a particular subset of carotenoid compounds, such as xanthophylls. Such compounds include but are not limited to the particular compounds described above, as well as, α -cryptoxanthin, β -cryptoxanthin, ζ -carotene, phytofluene, neurosporane, adonixanthin, echineneone, hydroxycanthaxanthin and the like. For a review of xanthophyll production, see Misawa, *et al.* (1995) *supra*). Using the methods of the invention, any compound of interest in the carotenoid pathway can be produced at high levels in a seed.

Secondary genes can also be selected to alter the fatty acid content of the plant for the production of specialty oils. For example, acyl-ACP thioesterase genes having specificity for particular fatty acid chain lengths may be used. See, for example, USPN 5,304,481, USPN 5,455,167, WO 95/13390, WO 94/10288, WO 92/20236, WO 91/16421, WO 97/12047 and WO 96/36719. Other fatty acid biosynthesis genes of interest include, but are not limited to, β-keto acyl-ACP synthases (USPN 5,510,255),

fatty acyl CoA synthases (USPN 5,455,947), fatty acyl reductases (USPN 5,370,996) and stearoyl-ACP desaturases (WO 91/13972).

Of particular interest is the use of a mangosteen acyl-ACP thioesterase as a secondary gene for fatty acid content modification. As described in WO 96/36719 and WO 97/12047, a high stearate content may be obtained in seeds by expression of a mangosteen acyl-ACP thioesterase. To combine the high oleic acid trait of the 3390 plants described herein with the 5266 high stearate plants described in WO 97/12047, crosses were made between 3390-1 and 5266-35 and between 3390-1 and 5266-5. Seeds resulting from these crosses contained oil having a high stearate, low linoleic, low linolenic and high carotenoid phenotype.

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Any means for producing a plant comprising the primary gene or both the primary and secondary genes are encompassed by the present invention. For example, the secondary gene of interest can be used to transform a plant at the same time as the primary gene either by inclusion of both expression constructs in a single transformation vector or by using separate vector, each of which express desired primary or secondary genes. The secondary gene can be introduced into a plant which has already been transformed with the primary gene, or alternatively, transformed plants, one expressing the primary gene and one expressing the secondary gene, can be crossed to bring the genes together in the same plant.

By combining the genes with tissue specific promoters, the carotenoid levels can be altered in particular tissues of the plant. Thus, carotenoid levels in the seed, including embryos and endosperm, can be altered by the use of seed specific transcriptional initiation regions. Such regions are disclosed, for example, in U.S. Patent No. 5,420,034, which disclosure is herein incorporated by reference.

In this manner, the transformed seed provides a factory for the production of modified oils. The modified oil may be used or alternatively, the compounds in the oils can be isolated. Thus, the present invention allows for the production of particular compounds of interest as well as speciality oils.

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The primary or secondary genes encoding the enzymes of interest can be used in expression cassettes for expression in the transformed plant tissues. To alter the carotenoid or fatty acid levels in a plant of interest, the plant is transformed with at least one expression cassette comprising a transcriptional initiation region linked to a gene of interest. Such an expression cassette is provided with a plurality of restriction sites for insertion of the gene of interest to be under the transcriptional regulation of the regulatory regions.

The transcriptional initiation may be native or analogous to the host or foreign or heterologous to the host. By foreign is intended that the transcriptional initiation region is not found the wild-type host into which the transcriptional initiation region is introduced.

Of particular interest are those transcriptional initiation regions associated with storage proteins, such as napin, cruciferin, \(\beta\)-conglycinin, phaseolin, or the like, and proteins involved in fatty acid biosynthesis, such as acyl carrier protein (ACP). See, U.S. Patent No. 5,420,034, herein incorporated by reference.

The transcriptional cassette will include the in 5'-3' direction of transcription, a transcriptional and translational initiation region, a DNA sequence of interest, and a transcriptional and translational termination region functional in plants. The termination region may be native with the transcriptional initiation region, may be native with the DNA sequence of interest, or may be derived from another source. Convenient termination regions are available from the Ti-plasmid of *A. tumefaciens*, such as the octopine synthase and nopaline synthase termination regions. See also, Guerineau et al., (1991), *Mol. Gen. Genet.*, 262:141-144; Proudfoot, (1991), *Cell*, 64:671-674; Sanfacon et al., (1991), *Genes Dev.*, 5:141-149; Mogen et al., (1990), *Plant Cell*, 2:1261-1272;

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Munroe et al., (1990), Gene, 91:151-158; Ballas et al., (1989), Nucleic Acids Res., 17:7891-7903; Joshi et al., (1987), Nucleic Acid Res., 15:9627-9639).

For the most part, the genes of interest of the present invention will be targeted to plastids, such as chloroplasts, for expression. Thus, the carotenoid biosynthesis gene or genes of interest may be inserted into the plastid for expression with appropriate plastid constructs and regulatory elements. Alternatively, nuclear transformation may be used in which case the expression cassette will contain a gene encoding a transit peptide to direct the carotenoid biosynthesis gene of interest to the plastid. Such transit peptides are known in the art. See, for example, Von Heijne et al. (1991) Plant Mol. Biol. Rep. 9:104-126; Clark et al. (1989) J. Biol. Chem. 264:17544-17550; della-Cioppa et al. (1987) Plant Physiol. 84:965-968; Romer et al. (1993) Biochem. Biophys. Res Commun. 196:1414-1421; and, Shah et al. (1986) Science 233:478-481. Plant carotenoid genes useful in the invention may utilize native or heterologous transit peptides.

It is noted that where the gene or DNA sequence of interest is an antisense DNA, targeting to a plastid is not required. In addition, where antisense inhibition of a given carotenoid biosynthesis gene is desired, the entire DNA sequence derived from the carotenoid biosynthesis gene is not required.

The construct may also include any other necessary regulators such as plant translational consensus sequences (Joshi, C.P., (1987), Nucleic Acids Research, 15:6643-6653), introns (Luehrsen and Walbot, (1991), Mol. Gen. Genet., 225:81-93) and the like, operably linked to the nucleotide sequence of interest.

It may be beneficial to include 5' leader sequences in the expression cassette which can act to enhance translation. Translation leaders are known in the art and include: picornavirus leaders, for example, EMCV leader (Encephalomyocarditis 5' noncoding region) (Elroy-Stein, O., Fuerst, T.R., and Moss, B. (1989) *PNAS USA* 86:6126-6130); potyvirus leaders, for example, TEV leader (Tobacco Etch Virus)

(Allison et al., (1986); MDMV leader (Maize Dwarf Mosaic Virus); Virology, 154:9-20), and human immunoglobulin heavy-chain binding protein (BiP), (Macejak, D.G., and Sarnow, P., (1991), Nature, 353:90-94; untranslated leader from the coat protein mRNA of alfalfa mosaic virus (AMV RNA 4), (Jobling, S.A., and Gehrke, L., (1987), Nature, 325:622-625; tobacco mosaic virus leader (TMV), (Gallie, D.R. et al., (1989), Molecular_Biology of RNA, pages 237-256; and maize chlorotic mottle virus leader (MCMV) (Lommel, S.A. et al., (1991), Virology, 81:382-385. See also, Della-Cioppa et al., (1987), Plant Physiology, 84:965-968.

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Depending upon where the DNA sequence of interest is to be expressed, it may be desirable to synthesize the sequence with plant preferred codons, or alternatively with chloroplast preferred codons. The plant preferred codons may be determined from the codons of highest frequency in the proteins expressed in the largest amount in the particular plant species of interest. See, EPA 0359472; EPA 0385962; WO 91/16432; Perlak et al. (1991) Proc. Natl. Acad. Sci. USA 88:3324-3328; and Murray et al. (1989) Nucleic Acids Research 17: 477-498. In this manner, the nucleotide sequences can be optimized for expression in any plant. It is recognized that all or any part of the gene sequence may be optimized or synthetic. That is, synthetic or partially optimized sequences may also be used. For the construction of chloroplast preferred genes, see USPN 5,545,817.

In preparing the transcription cassette, the various DNA fragments may be manipulated, so as to provide for the DNA sequences in the proper orientation and, as appropriate in the proper reading frame. Towards this end, adapters or linkers may be employed to join the DNA fragments or other manipulations may be involved to provide for convenient restriction sites, removal of superfluous DNA, removal of restriction sites, or the like. For this purpose, *in vitro* mutagenesis, primer repair, restriction,

annealing, resection, ligation, or the like may be employed, where insertions, deletions or substitutions, e.g. transitions and transversions, may be involved.

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The recombinant DNA molecules of the invention can be introduced into the plant cell in a number of art-recognized ways. Those skilled in the art will appreciate that the choice of method might depend on the type of plant, i.e. monocot or dicot, targeted for transformation. Suitable methods of transforming plant cells include microinjection (Crossway et al. (1986) BioTechniques 4:320-334), electroporation (Riggs et al. (1986) Proc. Natl. Acad. Sci. USA 83:5602-5606, Agrobacterium mediated transformation (Hinchee et al. (1988) Biotechnology 6:915-921) and ballistic particle acceleration (see, for example, Sanford et al., U.S. Patent 4,945,050; and McCabe et al. (1988) Biotechnology 6:923-926). Also see, Weissinger et al. (1988) Annual Rev. Genet. 22:421-477; Sanford et al. (1987) Particulate Science and Technology 5:27-37(onion); Christou et al. (1988) Plant Physiol. 87:671-674(soybean); McCabe et al. (1988) Bio/Technology 6:923-926 (soybean); Datta et al. (1990) Biotechnology 8:736-740(rice); Klein et al. (1988) Proc. Natl. Acad. Sci. USA, 85:4305-4309(maize); Klein et al. (1988) Biotechnology 6:559-563 (maize); Klein et al. (1988) Plant Physiol. 91:440-444(maize); Fromm et al. (1990) Biotechnology 8:833-839; and Gordon-Kamm et al. (1990) Plant Cell 2:603-618 (maize).

Alternatively, a plant plastid can be transformed directly. Stable transformation of chloroplasts has been reported in higher plants, see, for example, Svab et al. (1990) Proc. Nat'l. Acad. Sci. USA 87:8526-8530; Svab & Maliga (1993) Proc. Nat'l Acad. Sci. USA 90:913-917; Staub & Maliga (1993) Embo J. 12:601-606. The method relies on particle gun delivery of DNA containing a selectable marker and targeting of the DNA to the plastid genome through homologous recombination. In such methods, plastid gene expression can be accomplished by use of a plastid gene promoter or by transactivation of a silent plastid-borne transgene positioned for expression from a selective

promoter sequence such as that recognized by T7 RNA polymerase. The silent plastid gene is activated by expression of the specific RNA polymerase from a nuclear expression construct and targeting of the polymerase to the plastid by use of a transit peptide. Tissue-specific expression may be obtained in such a method by use of a nuclear-encoded and plastid-directed specific RNA polymerase expressed from a suitable plant tissue specific promoter. Such a system has been reported in McBride et al. (1994) Proc. Natl. Acad. Sci., USA 91:7301-7305.

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The cells which have been transformed may be grown into plants in accordance with conventional ways. See, for example, McCormick et al., Plant Cell Reports (1986), 5:81-84. These plants may then be grown, and either self or crossed with a different plant strain, and the resulting homozygotes or hybrids having the desired phenotypic characteristic identified. Two or more generations may be grown to ensure that the subject phenotypic characteristic is stably maintained and inherited and then seeds harvested to ensure the desired phenotype or other property has been achieved.

As a host cell, any plant variety may be employed. Of particular interest, are plant species which provide seeds of interest. For the most part, plants will be chosen where the seed is produced in high amounts, a seed-specific product of interest is involved, or the seed or a seed part is edible. Seeds of interest include the oil seeds, such as oilseed *Brassica* seeds, cotton seeds, soybean, safflower, sunflower, coconut, palm, and the like; grain seeds, *e.g.* wheat, barley, oats, amaranth, flax, rye, triticale, rice, corn, etc.; other edible seeds or seeds with edible parts including pumpkin, squash, sesame, poppy, grape, mung beans, peanut, peas, beans, radish, alfalfa, cocoa, coffee, tree nuts such as walnuts, almonds, pecans, chick-peas etc.

It is noted that the methods of the present invention have been demonstrated to provide increased carotenoid production in both oilseed *Brassica*, which has a green embryo, and in cotton, which has a white embryo.

In seed of cotton plants transformed with phytoene synthase, increases of total carotenoid levels ranging from 10 to 300 fold may be obtained. The majority of the increase in carotenoid levels, in this case, about 80%, is observed as an increase in phytoene levels. Increases in lutein levels are also obtained in this case, ranging from 1.5 to a 5 fold increase. In addition, α -carotene and β -carotene levels are also increased 10 to 100 fold, with β -carotene levels being 20 fold higher than α -carotene levels. Thus, as seen with Brassica, a second early carotenoid biosynthesis gene, such as phytoene desaturase, may be used with crtB to increase the metabolic flux through the carotenoid/isoprenoid pathway in cotton to produce a particular carotenoid.

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Furthermore, it should also be noted that the methods of the present invention have also been demonstrated herein to provide increased carotenoid production in additional plant species, such as *Arabidopsis* and corn.

In seed of Arabidopsis plants transformed with phytoene synthase, increases of total carotenoid levels ranging from 3 to at least approximately 20 fold may be obtained. A large increase in the level of β -carotene, ranging from 10 to 70 fold, are observed in seeds of transgenic Arabidopsis plants. Increases in lutein levels are also obtained in this case, ranging from 1.5 to a 3 fold increase. In addition, phytoene, α -carotene and lycopene levels are also increased. However, such increases in α -carotene, phytoene and lycopene are difficult to quantify as these levels are too low to measure in nontransformed control plants. Thus, as seen with Brassica and cotton, a second early carotenoid biosynthesis gene may be used with crtB to increase the metabolic flux through the carotenoid/ isoprenoid pathway in cotton to produce a particular carotenoid and to reduce the increased levels of phytoene.

In seed of corn plants transformed with phytoene synthase, increases of total carotenoid levels ranging from 2 to at least approximately 5 fold are obtained. The majority of the increase is seen in the levels of phytoene, while additional increases were

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observed in β-carotene levels. Increases of up to about 15 fold increases in β-carotene levels were obtained. Thus, again as with Brassica, cotton, and *Arabidopsis*, a second early carotenoid biosynthesis gene can be employed with the *crt*B to increase the metabolic flux through the carotenoid/isoprenoid pathway in corn to produce a particular carotenoid and to reduce the increased levels of phytoene. Additionally, additional genes, including secondary carotenoid biosynthesis genes can also be used to direct the production of particular carotenoids and xanthophyls.

In one embodiment of the invention, seed transcriptional initiation regions are used in combination with at least one carotenoid biosynthesis gene. This increases the activity of the carotenoid pathway and alters carotenoid levels in the transformed seed. In this manner, particular genes can be selected to promote the formation of compounds of interest. Where the gene selected is an early carotenoid biosynthesis gene the transformed seed has a significant increase in carotenoid biosynthesis as the result of an increase in the flux through the pathway. For Brassica seeds transformed with an early carotenoid biosynthesis gene, significant increases in the production of α -carotene, β -carotene and smaller increases in lutein in the seed oil, as well as altered oil fatty acid compositions are obtained. Seeds obtained from corn plants transformed with an early carotenoid biosynthesis gene also demonstrate an increased amount of carotenoid production.

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Where the early carotenoid biosynthesis gene is phytoene synthase, significant increases of a particular carotenoid include those ranging from a 10 to a 50 fold increase, preferably at least a 50 to a 100 fold increase, more preferably, at least a 50 to a 200 fold increase, such as the increases seen in α -carotene and β -carotene levels. Lutein levels, in this case, are also increased, but lower increases of 1.5 - 2 fold are obtained. At the same time, total carotenoid levels may be increased at least 10 to 25 fold, preferably 25 to 60 fold, and more preferably 25 to 100 fold. Thus, a seed of the invention

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transformed with a phytoene synthase gene has a substantial increase in levels of α - and β -carotene and total carotenoids, as well as smaller increases in lutein and other carotenoids, including phytoene. In some cases, it is not possible to quantitate the fold increase in a given carotenoid compound, as the levels are too low to detect in seeds from comparable non-transformed plants. In *Brassica napus*, for example, α -cryptoxanthin, lycopene, phytoene and phytofluene are all detected in various levels in seeds transformed with a *crt*B gene, but are not detectable in seeds from untransformed *Brassica napus* plants.

Where the early carotenoid biosynthesis gene is GGPP synthase or phytoene desaturase, 1.5 to 2 fold increases in lutein and \(\beta\)-carotene have been obtained in at least one transgenic plant for each gene. Lycopene is also detected in seeds from \(Brassica\) napus plants transformed with a \(crtE\) (GGPP synthase) gene. Total carotenoids in \(crtE\) or \(crtI\) transformants are also increased approximately 2 fold. Chlorophyll levels are also increased in \(B.\) napus transgenic plants expressing a \(crtE\) gene suggesting an increase in the levels of geranylgeranyl pyrophosphate (GGPP), which is the branch point substrate for carotenoid, chlorophyll and tocopherol biosynthesis. Increases in chlorophyll levels of 1.5 to 2 fold may be obtained in developing and mature seeds. Thus, also of interest as sources of carotenoids are plants which have been engineered to express increased levels of both \(crtB\) and \(crtE\).

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As demonstrated herein, the effect of one early carotenoid biosynthesis gene on the metabolic energy flux through the carotenoid pathway may be further effected by the addition of a second early carotenoid biosynthesis gene. Thus, the addition of a second early carotenoid biosynthesis gene for increasing the metabolic flow through the carotenoid biosynthesis pathway is also of interest in the present invention, and may find use for production of particular carotenoids either in the presence or absence of a secondary carotenoid biosynthesis gene.

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Where the early carotenoid biosynthesis gene phytoene synthase is cotransformed into *Brassica napus* with a second early carotenoid biosynthesis gene, phytoene desaturase, significant increases of particular carotenoids include increases in α-carotene, β-carotene, and lutein such as observed by expression of *crt*B alone. In addition, lycopene and phytoene levels are also increased in such plants, but increases are difficult to quantitate as these levels are too low to be detected in untransformed *Brassica napus* plants.

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Furthermore, when crtI and crtB are both expressed, total carotenoid levels greater than those observed with crtB alone may be obtained. In at least one plant, total carotenoid levels of 1.5 fold those observed in crtB plants were obtained. Lycopene levels are also increased over levels obtained in seeds of plants transformed with crtB alone. Lycopene levels may be increased from 4 to 15 fold over those obtained in seed of a homozygous crtB plant. In addition, a reduction in the ratio of phytoene to total carotenoids is also obtained, and as a result, levels of α -carotene and β -carotene are increased 1.2 to 1.8 fold over those obtained with crtB alone. In seeds of plants transformed with phytoene synthase alone, phytoene levels constituted as much as 20% of total carotenoids, while in plants cotransformed with phytoene synthase and phytoene desaturase, phytoene levels represent only 4% to 7% of the total carotenoids.

This metabolic energy effected by transformation with an early carotenoid gene can be funneled into a metabolic compound of choice by transformation with a second gene. As discussed above, the second gene is designed to promote the synthesis of a particular carotenoid by promoting the formation of the carotenoid of interest or alternatively by stopping the pathway to allow for the buildup of compounds. Therefore, significant amounts of carotenoids of interest can be produced in the transformed seeds of the present invention.

Where the primary carotenoid biosynthesis gene phytoene synthase is cotransformed with a secondary carotenoid biosynthesis gene, β -carotene ketolase, increases in levels of α -carotene, β -carotene and phytoene, such as those seen with transformation with crtB alone, are obtained. Furthermore, echinenone and canthaxanthin levels are also increased. However, such increases are difficult to quantitate as echinenone and canthaxanthin are either not produced in Brassica napus, or the levels are too low to be detected in B. napus plants expressing phytoene synthase alone and nontransformed control plants. Thus, for the production of a specific carotenoid, such as astaxanthin, the addition of a third carotenoid biosynthesis gene, such as β -carotene hydroxylase (crtZ), may find use in the present invention. Furthermore, the addition of a fourth carotenoid biosynthesis gene, such as phytoene desaturase, may also find use in the present invention.

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It should be noted that the carotenoid echinenone is a reaction intermediate in the production of canthoxanthin from β -carotene. The β -carotene ketolase (crtW) could react with the β -ring of α - or β -carotene. One β -ring reaction in β -carotene results in echinenone, two β -ring reactions in β -carotene form canthaxanthin, and one β -ring reaction in α -carotene makes 4-keto- α -carotene. This enzyme can not react with the ϵ -ring of α -carotene. Thus, two additional peaks on the HPLC chromatogram are produced in similar amounts, one representing echinenenone, and the other may represent 4-keto- α -carotene.

Where the primary carotenoid biosynthesis gene phytoene synthase is cotransformed with an antisense secondary carotenoid biosynthsis gene, ϵ -cyclase, large increases in levels of α -carotene, β -carotene and phytoene, such as those seen with transformation with crtB alone, are obtained. Some difference in the ratio of β -carotene to α -carotene is observed as compared to plants transformed with crtB alone, but large increases in both α -carotene and β -carotene levels are still observed. Lutein levels, on

the other hand, are either unchanged, increased, or in some cases decreased by as much as 80% as compared to seeds of untransformed control plants.

Initiation of carotenoid biosynthesis begins at approximately 15 days post anthesis in *B. napus* seeds, while expression of transformed genes utilizing the napin promoter begins about 18 days post anthesis. Thus, in order to more tightly control the α-carotene pathway to allow for the build up of β-carotene pathway carotenoids using antisense ε-cyclase, an earlier promoter, such as that of the *Lesquerella* kappa hyrodoxylase (described in pending U.S. patent application 08/898,038, filed 18 July, 1997), may find use. Thus, for increasing levels of a particular carotenoid using antisense, an earlier seed specific tanscriptional initiation region, may be used with a secondary carotenoid biosynthesis gene.

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The seeds of the invention which have been transformed with the primary early carotenoid biosynthesis gene also provide a source for novel oil compositions. The use of phytoene synthase as the primary gene, for example, results in substantial increases in oleic acid content in seed oil. By substantial increase is intended an increase of from about 5% to about 40%, specifically from about 20% to about 40%, more specifically from about 30% to about 40%. Thus, the seeds of the invention which have been transformed with a primary early carotenoid biosynthesis gene provide a source for modified oils having a high oleic acid content. That is, carotenoid biosynthesis genes, particularly early carotenoid biosynthesis genes can be used to produce seeds having at least 70% oleic acid, on a weight percentage basis. The oleic acid content in any seed can be altered by the present methods, even those seeds having a naturally high oleic acid contents. Alteration of seeds having naturally high oleic acid contents by the present methods can result in total oleic acid contents of as high as 80%.

Importantly, there is also a decrease in linoleic and linolenic acid content. By decrease in linoleic fatty acid content is intended a decrease from about 10% to about

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25%, preferably about 25% to about 40%, more preferably about 35% to about 60%. By decrease in linolenic fatty acid content is intended a decrease from about 10% to about 30%, preferably about 30% to about 60%, more preferably about 50% to about 75%. Thus, the methods of the invention result in oils which are more oxidatively stable than the naturally occurring oils. The modified oils of the invention are low-saturate, high oleic and low linolenic. Furthermore, the present invention provides oils high in monounsaturated fatty acids which are important as a dietary constituent.

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Based on the methods disclosed herein, seed oil can be modified to engineer an oil with a high oleic acid content as well as a high level of a carotenoid of interest. High oleic acid and high α - and β -carotene oils would have a longer shelf life as both the oleic acid and α - and β -carotene content would lend stability. It is also noted that such oils are more desirable as sources of carotenoids than the natural red palm oil, which oil contains high levels of saturated fatty acids.

The transformed seed of the invention can thus provide a source of carotenoid products as well as modified fatty acids. Where the intent is to produce particular carotenoid compounds of interest, methods are available in the art for the purification of the carotenoid compounds. In the same manner, methods available in the art can be utilized to produce oils purified of carotenoids. See, generally, WO 96/13149 and Favati et al. (1988) J. Food Sci. 53:1532 and the references cited therein.

That is, transformed seed and embryos can be visually determined and selected based on color as a result of the increased carotenoid content. The transformed seeds or embryos display a color ranging from yellow to orange to red as a result of the increased carotenoid levels. Therefore, where plant transformation methods involve an embryonic stage, such as in transformation of cotton or soybean, the carotenoid gene can be used in plant transformation experiments as a marker gene to allow for visual selection of

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transformants. Likewise, segregating seed can easily be identified as described further in the examples.

The following examples are offered by way of illustration and not by way of limitation.

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EXPERIMENTAL

Example 1 Expression Construct and Plant Transformation

A. SSU fusions to E. uredovora carotenoid biosynthesis genes

(1) Phytoene Synthase

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The SSU leader and crtB gene sequences were joined by PCR. The sequence of the SSU/crtB fusion is shown in Figure 1. The crtB gene from nucleotides 5057 to 5363 (numbering according to Misawa et al. (1990) supra) was joined to the SSU leader as follows. A Bg/II site was included upstream of the SSU leader start site to facilitate cloning. The thymidine nucleotide at 5057 of crtB was changed to an adenosine to make the first amino acid at the SSU leader/crtB junction a methionine, and the splice site a cys-met-asn. The native splice site for SSU is csy-met-gln. Note that Misawa et al. (1990) supra) indicates that the start site for the coding region for crtB is at nucleotide 5096. Thus, there are 13 amino acids upstream of the published start of the coding region for crtB and after the SSU splice site in the crtB/SSU fusion. Twelve of these amino acids are translated from Erwinia crtB upstream sequence and one is the added methionine. The crtB from 5363 (EcoRV) to 6009 (EcoRI) was then attached to the SSU-crtB fusion to obtain a complete SSU-crtB fusion construct designated pCGN3373 (Fig. 1).

25 (2) Phytoene Desaturase

A plasmid comprising a *E. uredovora crt*I gene fused to the transit peptide sequence of the pea Rubisco small subunit was described by Misawa *et al.* (*The Plant Journal* (1993) 4:833-840. An approximately 2.1 kb *Xbal/Eco*RI fragment of this plasmid containing the SSU-*crtI* fusion and a nos 3' termination region was cloned in position for expression from a napin 5' promoter.

(3) GGPP Synthase

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A similar construct containing the SSU transit fused to an *E. uredovora crt*E gene was obtained. The SSU-*crt*E fusion is present on an approximately 1.2 kb *BglII/Bam*HI fragment in pCGN3360.

B. SSU fusions to A. auriantiacum carotenoid biosynthesis genes

(4) beta-Carotene Hydroxylase (crtZ)

The SSU leader and crtZ gene sequences were joined by PCR. The crtZ gene (Misawa, et al. (1995) supra) nucleotide sequence was resynthesized to adjust for plant codon usage. The re synthesized crtZ gene was joined to the SSU leader by PCR as follows. A BglII site was included upstream of the SSU leader translation start site and a XhoI site was included downstream of the crtZ stop codon to facilitate cloning in the napin expression cassette. The nucleotide sequence of the complete ssu:crtZ fusion is shown in Figure 15.

(5) beta-Carotene Ketolase (crtW)

The SSU leader and crtW gene sequences were joined by PCR. The crtW gene (Misawa, et al. (1995) supra) nucleotide sequence was resynthesized to adjust for plant codon usage. The re synthesized crtW gene was joined to the SSU leader by PCR as follows. A BglII site was included upstream of the SSU leader translation start site and a XhoI site was included downstream of the stop codon to facilitate cloning in the napin expression cassette. The nucleotide sequence of the complete ssu:crtW fusion is shown in Figure 16.

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C. Expression Constructs for Plant Transformation

(1) Phytoene Synthase

pCGN3373 carrying the complete SSU/crtB fusion was cut with BglII and BamHI to excise the SSU/crtB fusion. The resulting fragment was ligated into the napin expression cassette in pCGN3223 at the BamHI site (see WO 94/10288 for description of napin expression cassette). The resulting construct, pCGN3389, was digested with

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HindIII to excise the napin 5'-SSU/crtB-napin 3' fragment, which was then cloned into HindIII cut pCGN1559PASS yielding pCGN3390. pCGN1559PASS is a binary vector for Agrobacterium-mediated transformation such as those described by McBride et al. (Plant Mol. Biol. (1990) 14:269-276) and is prepared from pCGN1559 by substitution of the pCGN1559 linker region with a linker region containing the following restriction digestion sites: Asp718/AscI/PacI/XbaI/BamHI/SwaI/Sse8387(PstI)/HindIII. A map of pCGN3390 is provided in Figure 2A. For expression of phytoene synthase in the corn endosperm, the crtB coding sequence from E. herbicola (Application WO 91/13078, Armstrong et al. (1990) supra) was cloned to be expressed under control from the rice glutelin, pGt1, promoter (Leisy, D.J. et al., Plant Mol. Biol. 14 (1989) 41-50) and the HSP70 intron sequence (U.S. Patent Number 5,593,874). This cassette also includes the transcriptional termination region downstream of the cloning site of nopaline synthase, nos 3' (Depicker et al., J. Molec. Appl. Genet. (1982) 1: 562-573) to create the vector pCGN9039 (Figure 2J) for transformation into corn.

The phytoene synthase coding sequence from corn (Figure 19) was also PCR amplified, fused with the SSU leader and cloned to be expressed from the napin promoter to create the expression construct pCGN9061.

(2) Phytoene Desaturase

A fragment comprising a napin 5'/SSU-crtI fusion/nos 3' construct as described above was cloned into a binary vector for plant transformation resulting in pCGN9010.

A map of pCGN9010 is provided in Figure 2C.

(3) GGPP Synthase

pCGN3360 carrying the complete SSU/crtE fusion was cut with BgIII and BamHI to excise the SSU/crtE fusion. The resulting 1.2 kb fragment was ligated into the napin expression cassette in pCGN3223 at the BamHI site. The resulting construct,

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pCGN3391, was digested with *HindIII* to excise the napin promoter-SSU/crtE napin 3' fragment, which was then cloned into *HindIII* cut pCGN1559PASS yielding pCGN3392. A map of pCGN3392 is provided in Figure 2B.

(4) Phytoene Synthase + Phytoene Desaturase

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The napin 5'-SSU/crtB-napin 3' fragment from pCGN3389 and the napin 5'/SSU-crtI fusion/nos 3' as present in pCGN9010 were inserted into a binary vector resulting in pCGN9009, shown in Figure 2D.

(5) Antisense Epsilon Cyclase + Phytoene Synthase

Brassica napus epsilon cyclase genes are isolated by PCR using primers designed from an Arabidopsis epsilon cyclase gene (Cunningham FX Jr (1996) Plant Cell 8:1613-1626). Sequence of B. napus epsilon cyclase genes is provided in Figures 9 (clone 9-4) and 10 (clone 7-6). An antisense construct is prepared by cloning anXhol/BamHI fragment of cDNA clone 9-4 into a napin expression cassette (pCGN3223) digested with Xhol and Bg/II. The napin 5'-antisense epsilon cyclasenapin 3' fragment is cloned along with a napin 5'-SSU/crtB-napin 3' fragment, fragment into a binary vector for plant transformation, resulting in pCGN9002, shown in Figure 2E.

(6) Antisense Beta Cyclase + Phytoene Synthase

Brassica napus beta cyclase genes are isolated by PCR using primers designed from an Arabidopsis beta cyclase gene (Cunningham FX Jr (1996) Plant Cell 8:1613-1626). Sequence of a B. napus beta cyclase cDNA, 32-3, is provided in Figures 11. An antisense construct is prepared by cloning anXhoI fragment of the beta cyclase cDNA clone into a napin expression cassette (pCGN3223) digested with XhoI. A clone containing the beta cyclase in the antisense orientation is selected. The napin 5'-antisense beta cyclase-napin 3' fragment is cloned along with a napin 5'-SSU/crtB-napin

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3' fragment into a binary vector for plant transformation, resulting in pCGN9017, shown in Figure 2F.

(7) beta-Carotene Hydroxylase + Phytoene synthase

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The vector pCGN9003 was constructed by removing the restriction sites between the crtB coding sequence and the napin 3' sequence by digestion with ClaI and XhoI and filling the ends with klenow creating the vector pCGN9000. PCGN9000 was digested with Asp718, and the fragment containing the napin 5'/SSU:crtB/napin 3' was ligated into the binary vector pCGN5139.

A binary vector for plant transformation, pCGN5139, was constructed using the neomycin phospho-transferase (nptII) kanamycin resistance gene driven by the CAMV 35S transcriptional initiation region (35S 5') and transcription termination (35S 3') sequences (Fraley et al., *Proc. Natl. Acad. Sci* (1983) 80:4803-4807, Gardner *et al.*, (1986) *Plant Mol Biol* 6:221-228). The 35S 5'-nptII-35S 3' fragment was then cloned into a vector containing ori322, Right border (0.5Kb), lacZ, Left Border (0.58Kb), as an *Xho* I fragment between the Right border-lacZ and Left border sequences. The ColEI and pRi origins of replication as well as the Gentamycin resistance gene were aquired from a derivative of pCGN1532 (McBride and Summerfelt, *Plant Molecular Biology*, (1990), 14:269-276). Finally, a linker containing unique restriction sites was synthesized and cloned into the *Asp* 718/ *Hind* III (within the lacZ sequence) sites to create the binary vector pCGN5139.

The plastid targeted ssu:crtZ fusion was cloned into the napin pCGN3223 seed expression cassette as a Bgl II -Xho I fragment to generate pCGN6203. The plasmid pCGN6203 carrying the complete napin cassette with ssu:crtZ was digested with NotI to excise the napin cassette containing the ssu:crtZ coding region. The excised fragment was ligated into the Not I site of the binary pCGN9003 carrying the napin SSU:crtB construct. The resulting construct, pCGN6205 (Figure 2H)

is a binary vector for Agrobacterium-mediated transformation such as those described by McBride et al. (Plant Mol. Biol. (1990) 14:269-276) and is prepared from pCGN1559 by substitution of the pCGN1559 linker region with a linker region containing the following restriction digestion sites: Asp718/AscI/PacI/XbaI/BamHI/SwaI/Sse8387(PstI)/HindIII. A map of pCGN6205 is provided in Figure 2H.

(8) beta-Carotene Ketolase + Phytoene synthase

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The ssu crtW plastid targeted fusion was cloned into the napin pCGN3223 seed expression cassette as a Bgl II -Xho I fragment to generate plasmid pCGN6202.

The plasmid pCGN6202 carrying the napin cassette with ssu:crtW was digested with NotI to excise a DNA fragment containing the napin cassette with ssu:crtZ. The resulting fragment was ligated into the Not I site of the binary pCGN9003 (described above) carrying the SSU:crtB napin construct. The resulting pCGN6204 (Figure 2G) is a binary vector for Agrobacterium-mediated transformation such as those described by McBride et al. (Plant Mol. Biol. (1990) 14:269-276) and is prepared from pCGN1559 by substitution of the pCGN1559 linker region with a linker region containing the following restriction digestion sites: Asp718/AscI/PacI/XbaI/
BamHI/SwaI/Sse8387(PstI)/HindIII. A map of pCGN6204 is provided in Figure 2G. (9) Phytoene synthase+ beta-Carotene hydroxylase+ beta-Carotene Ketolase

Construct pCGN6203 containing the napin cassette and ssu:crtZ was digested with Hind III to excise the fragment containing napin ssu:crtZ. The resulting HindIII fragment was cloned into the Hind III site of pCGN6204 to generate a triple crt genes binary pCGN6206 that contains napin ssu:crtB+ napin ssu:crtW+ napinssu:crtZ (Figure 2I).

D. Plant Transformation

Transformed Brassica napus plants containing the above described constructs are obtained as described in Radke et al. (Theor. Appl. Genet. (1988) 75:685-694 and Plant Cell Reports (1992) 11:499-505).

Transformed cotton plants, Gossypium hirsutum, containing phytoene synthase may be obtained using methods described in issued U.S. patent No. 5,004,863, and 5,159,135, and in Umbeck et al. (1987) Bio/Technology 5:263-266, or as described in copending application 08/539,176.

Transgenic Arabidopsis thaliana plants containing phytoene synthase may be obtained by Agrobacterium-mediated transformation as described by Valverkens et al., (Proc. Nat. Acad. Sci. (1988) 85:5536-5540), or as described by Bent et al. ((1994), Science 265:1856-1860), or Bechtold et al. ((1993), C.R.Acad.Sci, Life Sciences 316:1194-1199).

Microprojectile bombardment methods, such as described by Klein et al.

(Bio/Technology 10:286-291) may also be used to obtain nuclear transformed plants.

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Example 2 Analysis of Transgenic Plants

A. Visual Observations and Segregation Ratios

The napin-SSU leader/crtB plants in 212/86 were tagged at 21 days, 28 days and 35 days post anthesis. When the first plant, 3390-1 was harvested at 28 days, some of the seeds were obviously orange. AT 35dpa, the orange was obvious enough that a segregation ratio could be obtained. This trend of orange seeds has continued and is seen in each of the 17 lines harvested that have been obtained. A table of the segregation ratios is included below in Table 3.

TABLE 3

	Generation	Plant#	Orange	Green	Ratio	Chi Square
	T2	3390-1	291	88	3 to 1	0.64
5	T2	3390-2	150	22	No fit	
	T2	3390-8	293	87	3 to 1	0.90
	T2	3390-4	277	82	3 to 1	0.89
	T2	3390-5	243	62	3 to 1	1.90
	T2	3390-7	236	89	3 to 1	0.99
10	T2	3390-6	307	5	63 to 1	0.00
	T2	3390-3	121	50	No fit	1.64
	T2	3390-11	294	105	3 to 1	0.37 `
	T2	3390-15	287	83	3 to 1	1.30
	T2	3390-16	187	65	3 to 1	0.08
15	T2	3390-17	105	104	No fit	
	T2	3390-12	119	28	3 to 1	2.78
	T2	3390-14	283	107	3 to 1	1.23
	T2	3390-19	238	94	3 to 1	1.94
	T2	3390-20	251	4	63 to 1	0.00
20	T2	3390-27	229	4	63 to 1	0.04

B. Carotenoid Analysis of Developing Seeds

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Carotenoids were extracted from seeds harvested at approximately 35 days post-anthesis as follows. Eight seed samples of orange seeds from transgenic plant 3390-1 and eight seed samples of a 212/86 variety rapeseed control plant were ground in 200µl of 70% acetone/30% methanol. The ground seed mixture was then spun in a microcentrifuge for approximately 5 minutes and the supernatant removed. Two additional 70% acetone/30% methanol extractions were conducted with the pelleted seed material and all three supernatants pooled and labeled A/M extract.

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At this point in the extraction, the control seed pellets are white, whereas the seed pellets from the transgenic seeds have a yellow color. The pellets are then extracted twice with ether and the resultant supernatants pooled and labeled E extract. The A/M extract was then transferred to ether as follows. 450µl ether and 600µl of water were added to the extracts, followed by removal of the ether layers. The A/M extracts were then washed two more time with 400µl of ether, and the ether fractions from the three A/M washes pooled. The E extracts described above were washed once with 400µl of water and pooled with the A/M ether fractions. The pooled ether fractions were blown down to a volume of approximately 300µl with nitrogen gas and filtered using a syringe microfilter. The sample vials were rinsed with approximately 100µl ether and the rinse was similarly filtered and pooled with the initial filtrate, yielding total volume of approximately 150µl. A 50µl aliquot was stored at -20YC until further analysis and the remaining 100µl sample was saponified as follows. 100µl of 10% potassium hydroxide (KOH) in methanol was added to each 100µl sample and the mixture stored in the dark at room temperature for approximately 2 hours. 400µl of water was then added to the samples and the ether phase removed. For better phase separation, saturated NaCl may be substituted for the water. The water solution was then extracted twice more with 100µl of ether and the ether samples pooled and washed with water.

The saponified samples were then analyzed by HPLC analysis on a Rainin microsorb C18 column (25cm length, 4.6mm outside diameter) at a flow rate of 1.5ml per minute. The gradient used for elution is as follows:

A = acetonitrile

B = hexane/methylene chloride (1:1)

C = methanol.

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The initial solution was 70:20:10 (A:B:C). At 2.5 minutes the solution is ramped over 5 minutes to 65:25:10 (A:B:C) and held at this for 12.5 minutes. The solution is then ramped to 70:20:10 (A:B:C) over two minutes followed by a three minute delay prior to injection of the next sample. The absorbance of the eluting samples is continuously monitored at 450 and 280 nm and known chemical and biological standards were used to identify the various absorbance peaks.

In Figures 3 and 4, results of analyses of saponified samples are provided for control and pCGN3390 transformed seeds, respectively. Clear increases in the levels of α - and β -carotene and phytoene in the transagenic plant seeds are observed, as well as smaller increases in levels of the hydroxylated carotenoid, lutein.

C. Carotenoid and Tocopherol Analysis of Mature Seeds from crtB Transgenic Plants

Mature 3390 T2 seed were sent to an analytical laboratory for quantitative analysis using standard HPLC methods known in the art. These results of these analysis are shown in Table 4 below. Compound levels are presented as $\mu g/g$.

Seeds designated "Maroon" were selected based on seed color. The seeds which have orange embryos appear maroon colored at maturity as opposed to the black-brown appearance of seeds from wild type plants of this cultivar. Seeds designated as "Random" were not selected for color. As 3390-1 is segregating 3 to 1 for Kan, the "Random" population includes a proportion of nulls. The maroon population contains only transgenics. Due to an effort to exclude nulls from this population, the inclusion of homozygotes may be favored.

TABLE 4

	COMPOUND	CONTROL	3390-1 RANDOM	3390-1 <u>MAROON</u>
5	Lutein	7.2	18	26
	Zeaxanthin	nd*	nd	nd
	α-cryptoxanthin	nd	8	15
	β-cryptoxanthin	nd	nd	. nd
	Lycopene	nd	2.3	5.1
10	cis-Lycopene	nd	2.9	5.4
	α-carotene	0.6	124	244
	β-carotene	0.9	177	338
	cis-B-carotene	0.2	. 12	26
	Other	6	34	51
15	Total colored carotenoids	14.9	378.2	710.5
	Phytoene	\mathbf{nd}	62	139
	Phytofluene	nd	24	54
	Total all carotenoids	14.9	464.2	903.5
20	Alpha-tocopherol	74	93	109
	Gamma-tocopherol	246	188	95
	Delta-tocopherol	3	. 5	5

*nd = not detected

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In the non-transgenic sample, "other" includes mostly very polar compounds, such as neoxanthin, violaxanthin, etc. In the transgenic sample "other" includes these and additional compounds, such as zeta-carotene, neurosporene, and mono-cyclic carotenoids.

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Results of carotenoid analysis of 3390 T2 seeds from transformed plants of B. napus variety Quantum (SP30021) are presented in Figure 12.

Results of carotenoid analysis of 3390 T3 seeds from transformed plants of B.

napus variety 212/86 (SP001) are presented in Figure 13.

The above results demonstrate that α - and β -carotenes levels are significantly increased in the mature seeds as the result of expression of the crtB gene. Generally, the overall increase in carotenoids is quite high, nearly 50 fold for colored carotenoids and up to 60 fold if phytoene and phytofluene are included. It is clear that the flux through the isoprenoid pathway has been dramatically increased. Additionally it is noted that the α -tocopherol (Vitamin E) levels are also increased by nearly 50%.

D. Germination Studies

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Ten mature seeds of 3390-1 and 10 seeds of 212/86 control were planted in soil and grown in a walk-in growth chamber. The transgenics emerged 1 to 2 days later than the controls, however, all 10 seeds did germinate. The transgenics were yellowish-pink when they first emerged but greened up in one to two days. At the emergence of the first true leaf, no difference in color was observed. Plants germinated from both the transgenic and control seeds developed normally.

E. Fatty Acid Analysis

Fatty acid composition of mature seeds was determined by GC analysis of single T2 seeds harvested from trangenic plants 3390-1 and 3390-8. Single seeds from both Random (R) and Maroon (M) populations (as defined above) were analyzed and compared to seeds from a 212/86 control (SP001-1). The results of these analyses are provided in Table 5 below as weight % total fatty acids.

TABLE 5
FATTY ACID COMPOSITION OF 3390-1 AND 3390-8 LINES

SAMPLE	10:0	12:0	14:0	16:0	16:1	18:0	18:1	18:2	18:3	20:0	20:1	20:2	22:0
CONTROL	1.5	0	0.1	5.1	0.4	1.7	59.9	17.1	12.0	0.6	1.2	0.1	0.3
CONTROL	1.8	0	0.1	5.1	0.4	1.7	60.1	16.6	12.1	0.6	. 1.2	0.1	0.3
CONTROL	2.0	0	0.1	5.0	0.4	1.6	60.5	16.2	12.0	0.6	1.2	0.1	0.3
CONTROL	2.2	0	0.1	5.2	0.4	1.6	57.2	18.2	12.7	0.6	1.3	0.1	0.4
CONTROL	1.6	0	0.1	4.7	0.4	1.8	62.7	15.3	11.3	0.6	1.2	0.1	0.3
3390-1-R	2.8	0	0.1	4.8	0.5	3.6	69.9	10.6	4.8	1.2	1.1	0.0	0.6
3390-1-R*	1.5	0	0.1	4.7	0.3	1.5	58.1	19.3	12.3	0.5	1.2	0.1	0.3
3390-1-R	3.5	0	0.1	4.2	0.3	2.6	71.1	9.6	5.8	1.0	1.2	0.0	0.6
3390-1-R*	1.5	0	0.1	4.7	0.3	1.9	61.0	17.8	10.4	0.7	1.3	0.1	0.3
3390-1-R	2.2	0	0.1	4.4	0.3	3.1	73.6	8.9	4.4	1.2	1.1	0.0	0.7
3390-1-R	1.9	0	0.1	4.5	0.3	2.4	72.7	10.6	4.7	0.9	1.3	0.1	0.6
3390-1-R	2.5	0	0.1	4.2	0.3	3.4	71.7	10.0	5.1	1.1	1.0	0.0	0.6
3390-1-R	1.7	0	0.1	4.4	0.3	2.6	73.5	10.0	4.5	1.0	1.2	0.1	0.6
3390-1-R	1.9	0	0.1	4.2	0.3	2.3	72.4	9.9	6.3	0.9	1.2	0.1	0.5
3390-1-R	2.5	0	0.1	4.2	0.3	2.7	72.0	10.1	5.1	1.0	1.2	0.1	0.6
3390-1-R*	1.5	0	0.1	4.7	0.3	1.7	58.5	18.5	12.6	0.6	1.2	0.1	0.3
3390-1-R	2.8	0	0.1	4.6	0.4	3.7	71.8	9.1	4.2	1.3	1.2	0.0	0.7
3390-1-R	1.8	0	0.1	4.0	0.3	2.3	72.4	11.1	5.2	0.9	1.3	0.1	0.5
3390-1-R	1.7	0	0.1	4.4	0.3	2.7	73.9	9.9	4.2	1.0	1.2	0.1	0.6
3390-1-R	1.7	0	0.1	4.6	0.4	2.6	71.4	10.9	5.5	1.0	1.3	0.1	0.6
3390-1-R	2.7	0	0.1	4.2	0.3	2.8	72.1	9.9	5.0	1.1	1.3	0.0	0.6
3390-1-R	2.0	0	0.1	4.5	0.3	3.0	72.5	9.7	4.6	1.2	1.3	0.1	0.7
3390-1-R	1.8	0	0.1	4.9	0.4	3.4	71.8	10.4	4.2	1.2	1.2	0.0	0.7
3390-1-R*	0.9	0	0.1	4.5	0.3	1.7	55.9	18.8	15.6	0.5	1.3	0.1	0.3
3390-1-R*	1.4	0	0.1	4.8	0.4	1.7	57.1	18.0	14.4	0.6	1.2	0.1	0.3
3390-1-R*	1.4	0	0.1	4.5	0.3	1.7	57.8	18.5	13.5	0.6	1.3	0.1	0.3
3390-1-R	2.2	0	0.1	4.5	0.3	2.5	73.4	9.7	4.6	0.9	1.2	0.0	0.5
3390-1-R	1.5	0	0.1	3.8	0.3	2.7	75.9	8.1	4.6	1.0	1.4	0.0	0.6
3390-1-R	1.6	.0	0.1	4.5	0.3	2.6	71.9	10.6	5.5	1.0	1.3	0.1	0.6
3390-1-R*	1.3	0	0.1	6.2	0.5	1.4	53.6	21.7	13.2	0.5	1.1	0.1	0.3
3390-1-R	2.1	0	0.1	4.3	0.3	2.4	72.3	10.7	5.1	0.9	1.2	0.0	0.6
3390-1-R*	1.3	0	0.1	5.0	0.3	1.6	57.8	18.8	13.0	0.5	1.3	0.1	0.3
3390-1-R	2.1	0	0.1	4.4	0.3	3.3	72.7	9.2	4.8	1.2	1.2	0.0	0.7
3390-1-R	1.5	0	0.1	4.5	0.3	3.3	72.6	10.1	4.6	1.2	1.1	0.1	0.7
3390-1-R*	1.2	0	0.1	4.7	0.3	1.9	59.9	17.1	12.6	0.6	1.3	0.1	0.4
3390-1-M	2.8	0	0.1	4.0	0.3	2.8	69.8	10.6	7.1	0.9	1.2	0.0	0.4
3390-1-M	2.0	0	0.1	4.9	0.4	3.3	70.3	11.1	4.9	1.2	1.2	0.1	0.7
3390-1-M	1.5	0	0.1	4.4	0.3	3.2	73.4	9.5	4.3	1.3	1.3	0.0	8.0
3390-1-M	1.5	0	0.1	4.5	0.3	2.8	72.7	10.0	5.1	1.1	1.3	0.0	0.7
3390-1-M	1.8	0	0.1	4.2	0.3	3.1	73.5	9.6	4.7	1.1	1.2	0.0	0.6
3390-1-M	1.5	0	0.1	4.7	0.3	2.9	71.6	10.7	5.1	1.1	1.2	0.1	0.7
3390-1-M	1.5	0	0.1	4.5	0.3	3.2	72.6	10.2	4.3	1.2	1.3	0.0	0.8
3390-1-M	1.8	0	0.1	4.4	0.3	2.9	72.0	10.4	5.2	1.1	1.2	0.1	0.6
3390-1-M	1.5	0	0.1	4.4	0.3	2.6	73.6	10.0	4.5	1.1	1.2	0.1	0.7
3390-1-M	2.3	0	0.1	4.3	0.3	3.0	73.0	9.7	4.5	1.1	1.2	0.0	0.6

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SAMPLE	10:0	12:0	14:0	16:0	16:1	18:0	18:1	18:2	18:3	20:0	20:1	20:2	22:0
3390-8-R*	1.0	0	0.1	4.9	0.3	1.6	59.2	18.9	11.9	0.5	1.2	0.1	0.3
3390-8-R	2.1	0	0.1	4.2	0.3	2.7	71.9	10.2	5.6	1.0	1.2	0.1	0.6
3390-8-R	1.5	0	0.1	4.4	0.3	2.3	72.5	10.4	5.7	0.9	1.4	0.1	0.6
3390-8-R*	1.2	0	0.1	4.9	0.3	1.7	59.7	18.2	11.6	0.6	1.3	0.1	0.4
3390-8-R*	1.5	0	0.1	4.7	0.3	1.6	58.7	18.5	12.2	0.6	1.3	0.1	0.4
3390-8-R	1.8	0	0.1	4.2	0.3	2.9	73.4	9.2	5.2	1.1	1.3	0.0	0.6
3390-8-R*	1.1	0	0.1	4.7	0.3	1.5	56.9	19.3	14.1	0.5	1.1	0.1	0.2
3390-8-R	2.2	0	0.1	4.6	0.3	3.0	71.4	10.0	5.2	1.1	1.2	0.1	0.7
3390-8-R	1.7	0	0.1	4.6	0.4	2.4	72.5	11.0	4.8	0.9	1.3	0.1	0.5
3390-8-R	2.4	0	0.1	4.7	0.3	2.9	74.0	8.4	4.0	1.1	1.2	0.0	0.7
3390-8-R	1.9	0	0.1	4.6	0.4	3.0	72.7	9.7	4.8	1.0	1.2	0.0	0.6
3390-8-R	2.0	0	0.1	4.4	0.3	2.8	73.2	9.7	4.5	1.0	1.3	0.0	0.6
3390-8-R	1.5	0	0.1	4.3	0.3	2.6	71.8	10.7	5.8	1.0	1.3	0.1	0.6
3390-8-R	1.5	0	0.1	4.4	0.3	2.7	72.6	10.5	4.9	1.0	1.3	0.1	0.6
3390-8-R	2.0	0	0.1	4.9	0.4	3.3	71.1	10.4	4.9	1.1	1.1	0.1	0.6
3390-8-R	2.1.	0	0.0	4.5	0.4	3.6	73.0	8.8	4.3	1.3	1.2	0.0	0.7
3390-8-R	2.2	0	0.1	5.1	0.4	2.9	67.6	12.3	6.5	1.1	1.2	0.1	0.7
3390-8-R	1.8	0	0.1.	4.2	0.3	2.6	73.5	9.9	4.8	1.0	1.3	0.1	0.6
3390-8-R	1.7	0	0.1	4.7	0.3	3.0	72.5	9.9	4.6	1.2	1.3	0.1	0.7
3390-8-R	1.7	0	0.1	4.6	0.4	2.8	73.7	9.5	4.1	1.1	1.3	0.1	0.7
3390-8-R	1.5	0	0.1	4.5	0.3	3.0	74.7	8.5	4.2	1.2	1.2	0.0	0.7
3390-8-R	1.5	0	0.1	4.4	0.4	1.9	70.0	11.8	7.2	0.8	1.4	0.1	0.5
3390-8-R	1.7	0	0.1	4.4	0.3	2.5	71.8	11.1	5.2	1.0	1.3	0.1	0.6
3390-8-R	1.4	0	0.1	4.5	0.4	2.8	73.3	9.7	4.9	1.1	1.2	0.1	0.6
3390-8-R	1.5	0	0.1	4.8	0.4	3.0	72.6	10.6	4.1	1.1	1.2	0.1	0.7
3390-8-R*	1.4	0	0.1	5.8	0.4	2.9	54.0	20.0	13.0	0.8	1.1	0.1	0.4
3390-8-R	1.4	0	0.1	4.4	0.3	2.7	71.2	10.8	6.0	1.0	1.3	0.1	0.6
3390-8-R	1.7	0	0.1	4.6	0.4	2.8	72.6	10.0	5.1	1.0	1.2	0.1	0.6
3390-8-R*	1.0	0	0.1	4.6	0.3	1.6	59.6	18.5	12.3	0.5	1.2	0.1	0.3
3390-8-R*	1.1	0	0.1	4.6	0.3	1.4	56.5	20.4	13.4	0.5	1.3	0.1	0.3
3390-8-M	1.8	0	0.1	4.7	0.4	3.3	70.1	11.1	5.5	1.2	1.1	0.1	0.7
3390-8-M	1.5	0	0.1	4.3	0.3	3.0	73.0	10.3	4.3	1.1	1.2	0.1	0.7
3390-8 - M	1.9	0	0.1	4.5	0.4	3.7	73.1	8.9	4.2	1.3	1.2	0.0	0.7
3390-8-M	1.6	0	0.1	4.4	0.3	2.5	73.4	9.7	5.1	1.0	1.3	0.1	0.7
3390-8-M	1.3	0	0.1	4.4	0.3	3.0	73.7	9.6	4.4	1.1	1.3	. 0.0	0.7
3390-8-M	2.1	0	0.1	4.3	0.3	3.2	74.0	8.9	4.1	1.2	1.2	0.1	0.6
3390-8-M	2.1	0	0.1	3.9	0.3	1.6	71.6	11.9	5.7	0.7	1.5	0.1	0.5
3390-8-M	1.6	0	0.1	4.6	0.3	2.8	71.0	11.8	4.8	1.0	1.3	0.1	0.6
3390-8-M	2.1	0	0.1	4.8	0.4	3.2	70.3	10.7	5.2	1.2	1.2	0.1	0.7
3390-8-M	1.6	0	0.1	4.5	0.3	2.9	72.7	9.9	4.8	1.1	1.3	0.0	0.7

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The above data demonstrate a substantial increase in oleic acid (18:1) in seeds from each of the transgenic lines. The increase in oleic acid is at the expense of linoleic and linolenic acids, both of which were decreased in the transgenic lines. Increases in 18:0 and 20:0 fatty acids were also observed. Based on these data, the null seeds present in the Random population can be identified, and are marked on Table 5 with an asterisk (*). All of the seeds in the Maroon populations from each transgenic line have the observed altered fatty acid composition, confirming that the altered fatty acid composition is the result of expression of the *crt*B gene.

The trends in fatty acid composition data in the transgenic seeds which indicate positive and negative correlations of fatty acid composition changes with the observed increase in 18:1 levels are provided in Figures 5-7. The increase in 18:1 correlates with the decreases in 18:2 and 18:3. (Figure 5). The increase in 18:1 also correlates with an increase in both 18:0 and 20:0, but little effect on 16:0 was seen (Figure 6). The increase in 18:0 also correlated with an increase in 20:0 (Figure 7).

F. Carotenoid Analysis of Mature Seeds from crtE Transgenic Plants

Carotenoids were analyzed in mature T2 seeds of 3392 *B. napus* plants tranformed to express the *E. uredovora crt*E gene. Approximately two fold increases in levels of lutein and \(\beta\)-carotene was observed in seeds of plant 3392-SP30021-16. Lycopene was also detected in these seeds and is undetectable in seeds of untransformed control plants. Analysis of seeds from 7 additional 3392 transformants did not reveal significant increases in the carotenoid levels.

G. Analysis of Chlorophyll and Tocopherol Levels in crtE Transgenic Plants

Chlorophyll levels were analyzed using a spectrophotometric assay (Bruinsma, J. 1961, A comment on the spectrophotometric determination of chlororphyll, Biochem Biophy Acta, 52:576-578) in mature T2 seeds of transgenic 3392 *B. napus* plants.

Levels in 3392 transgenic plants were compared to seeds of transgenic *B. napus* plants

expressing phytoene synthase (crtB) and to nontransformed control plants. Results are shown in Table 6 below.

TABLE 6

Pigment concentration (μg/gFW)

Gene and sample	Total carotenoids	Total chlorophyll	
_			
Phytoene synthase		•	
27 DPA SP001 control	53	676	•
27 DPA T4 3390-1-6	354	. 282	
40 DPA SP001 control	47	471	
40 DPA T4 3390-1-6	534	179	
50 DPA SP001 control	16	125	
50 DPA T4 3390-1-6	648	125	
GGPP synthase			
35 DPA SP30021 control		68	407
35 DPA T2 3392-4		65	660
35 DPA T2 3392-16		73	648
Mature SP30021 control		21	35
Mature T2 3392-4		25	31
Mature T2 3392-16		50	60

Chlorophyll concentrations of the 35 DPA seeds of two lines were increased by approximately 60% compared to the levels of the control plant. The initial results demonstrate that the GGPP synthase gene increased the GGPP substrate availability for chlorophyll biosynthesis during seed development. Mature seeds of the 3392-16 line had higher chlorophyll and carotenoid concentrations than those of the control.

H. Carotenoid Analysis of Mature Seeds from crt I Transgenic Plants.

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Carotenoids were analyzed in mature T2 seeds of 9010 *B. napus* plants tranformed to express the antisense lycopene ε-cyclase gene. Seeds of nine transgenic plants were analyzed for carotenoid content. An approximately two fold increase in levels of lutein, β-carotene and total carotenoids was observed in seeds of one line, 9010-SP30021-10, when compared to control plants.

I. Carotenoid Analysis of Mature Seeds from crtB + crtI Transgenic Plants

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Carotenoid levels of Mature 9009 T2 seeds were extracted and quantified on an HPLC as follows. Approximately 100mg of seeds were ground in a mortar and pestle in 3ml extraction solvent (hexane/acetone/ethanol (50/25/25 v/v) with 0.2ml of an internal standard (5mg/ml β-apo-8' carotenal (dissolved in 100μl hexane), in acetonitrile/methylene chloride/methanol (50/40/10, v/v/)). The extraction solution was transferred to a new glass tube, and the remaining seed was again extracted with the extraction solvent and pooled with first extraction solution. The extraction was repeated until no color was visible in the extraction solution. Pooled extracts were mixed by vortexing briefly, then centrifuged for approximately 5 minutes. The resulting supernatant was transfered to a new tube and dried under nitrogen gas. The residue was resuspended in 2ml of hexane. Potassium hydroxide, in methanol, was added to a final concentration of 5%, and the solution was incubated overnight in the dark at 4°C. Another 2ml of hexane was then added to the solution with 1ml of saturated sodium chloride. The solution was mixed briefly by vortexing and centrifuged for approximately 5 minutes. The upper hexane layer was removed and transferred to a new glass tube. The remaining bottom phase was again extracted with hexane and centrifuged. The upper phase was combined with the previous hexane phase. This was repeated until the hexane phase was colorless. The pooled hexane phases were dried under nitrogen gas, and the residue was dissolved in 2.0ml of acetonitrile/methylene chloride/methanol (50/40/10 v/v). The solution was filtered through a 0.45µm filter and colected in a brown autopsampler vial. Carotenoid concentrations were determined on a Hewlett Packard 1050 High-Performance Liquid Chromatograph (HPLC), and isocratic separation of carotenoids was performed on a Hewlett Packard reverse phase C-18 (5µ) column (4.6 mm x 20cm) at 30°C. The mobile phase was acetonitrile/ methylene chloride/ methanol (80/10/10, v/v) with a flow rate of 1.0ml/min and a sample injection

volume of $20\mu l$ (running time of 22min). Routine detection of colored carotenoids is at 450 nm, phytoene at 280 nm, and phytofluene at 365 nm. Spectral scans for peak purity were made at 250 nm and 600 nm. Spectra of peaks at the upslope, apex, and downslope are normalized and overlaid. Superimposing spectra were taken as evidence of peak purity. The results are shown in Table 7 below. Carotenoid levels are presented as $\mu g/gFW$.

TABLE 7

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Sample ID #	Lutein	Lycopene	α-Carot	ene β-C	Carotene	Phyt	oene Total
_							
SP30021 control	36	ND	ND		4	ND	40
3390-SP001-1-6-15	5						
(T5 Homo)	54	4	552	638		277	1525
9009-SP30021-1	44	44	336	691		42	1157
9009-SP30021-6	53	87	689	1118		152	2099
.9009-SP30021-9	48	34	487	798		194	1561
9009-SP30021-10	33	25	248	489		34	829
9009-SP30021-12	31	ND	ND	2		ND	33
9009-SP30021-14	42	37	404	791		81	1355
9009-SP30021-15	37	15	137	278		ND	467
9009-SP30021-16	50	38	428	828		65	1409

The results demonstrate that as with plants transformed to express crtB alone, plants expressing crtB and crtI contain significant increases in total carotenoid levels. Furthermore, it is apparent that expression of crtI with crtB, leads to further modification of the phytoene pools which accumulate in crtB transformants. Phytoene levels were reduced from about 20% of total carotenoids in lines transformed with crtB alone, to 4% to 7% of total carotenoids in the crtB + crtI lines. This indicates that phytoene desaturase can have a synergistic effect with phytoene synthase in increasing the metabolic flux through the carotenoid/ isoprenoid pathway, and provides for even

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greater increases in a desired carotenoid compound, such as α -carotene and β -carotene, than is obtained by expression of crtB alone. The increased flux also appears to result in increased total carotenoid production, in addition to the composition shift from phytoene. For example, the carotenoid levels in the segregating T2 seed populations of 9009-10 are significantly higher than those detected in the 3390 homozygous seed population in 3390-1-6-15.

J. Carotenoid Analysis of mature Seeds from crtB + Antisense ε-Cyclase Transgenic Plants

Carotenoids from mature seeds from 9002 transformants were extracted and analyzed using the method described in example 2I above. These results are shown in Figure 14.

The initial results show a modification to the ratio of β -carotene to α -carotene. In addition, several lines show a significant reduction in lutein levels when compared to nontransgenic controls. In 9002 T2 lines, β -carotene to α -carotene ratios averaged 1.5, ranging from 1.1 to 2.5. For comparison, T2 3390 lines containing crtB, the ratio of β -carotene to α -carotene averaged 1.9, ranging from 1.5 to 2.4.

K. Carotenoid Analysis of Mature Seeds from crtB Transgenic Cotton Plants

Mature 3390 T2 seeds from cotton were collected and carotenoid extracts were prepared and analyzed according to the method described in 2I above. These results are shown in Table 8 below. Carotenoid levels are presented as µg/gFW.

TABLE 8

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Sample ID#	Lutein	Lycopene	α-Carotene	β-Carotene	Phytoene	Total
C130 control	2	ND	ND	ND	ND	2
3390-C130-5-1	7	ND	486	420	517	

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An approximately 3 fold increase in lutein was observed in seeds of plant 3390-C130-5-1. Alpha-carotene, β -carotene and phytoene were also observed in this line and are undetectable in nontransformed control plants. With β -carotenoid levels being 20 fold higher than those of α -carotene. Total carotenoid levels were increased by more than 250 fold, with phytoene accounting for approximately 80% of that total.

L. Carotenoid Analysis of Mature Seed from crtB + crtW Transgenic Plants

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Carotenoid levels of Mature 6204 T2 seeds were extracted and quantified on an HPLC as follows. Approximately 100mg of seeds were ground in a mortar and pestle in 3ml extraction solvent (hexane/acetone/ethanol (50/25/25 v/v) with 0.3ml of an internal standard (5mg/ml β-apo-8' carotenal (dissolved in 100μl hexane), in acetonitrile/methylene chloride/methanol (50/40/10, v/v/)). The extraction solution was transferred to a new glass tube, and the remaining seed was again extracted with the 2 ml extraction solvent and pooled with first extraction solution. The extraction was repeated until no color was visible in the extraction solution. Pooled extracts were mixed by vortexing briefly, then centrifuged for approximately 5 minutes. The resulting supernatant was transfered to a new tube and dried under nitrogen gas. The dried sample was stored in the dark overnight at 4°C. The residue was resuspended in 3ml of hexane and 1 ml methanol, and 1 ml of saturated sodium chloride was added and mixed. The samples were centrifuged briefly, and the upper phase was transferred to a new tube. The remaining bottom phase was again extracted with 2 ml hexane and centrifuged. The upper phase was combined with the previous hexane phase. This was repeated until the hexane phase was colorless. The pooled hexane phases were dried under nitrogen gas, and the residue was dissolved in 2.0ml of acetonitrile/methylene chloride/methanol (50/40/10 v/v). The solution was filtered through a 0.45µm filter and colected in a brown autopsampler vial. Carotenoid concentrations were determined on a Hewlett Packard 1100 High-Performance Liquid Chromatograph (HPLC), and isocratic separation of carotenoids was performed on a Spherisorb ODS2 reverse phase C-18 (5μ) column (4.6 mm x 25cm) at 30°C. The mobile phase was 82 acetonitrile/ 10 dioxane /8 methanol (v/v) containing 150 mM ammonium acetate/ 0.1 triethylamine, with a flow rate of 1.0ml/min and a sample injection volume of 20µl (running time of 46 min). Routine detection of colored carotenoids is at 450 nm, phytoene at 280 nm, and phytofluene at 365 nm. Spectral scans for peak purity were made at 250 nm and 600 nm. Spectra of peaks at the upslope, apex, and downslope are normalized and overlaid. Superimposing spectra were taken as evidence of peak purity. The results are shown in Table 10 below, and an HPLC chromatogram is shown in Figure 17. Table 9 below describes the relevant peak retention times shown in Figure 17. Carotenoid levels are presented as $\mu g/gFW$.

Table 9.

Ret Time	Area	Amt/Area	Amount	Compound
[min]	[mAU*s]		[ug/gFW]	Name
3.500			-	Astaxanthin
5.428	721.34	4.3×10^{-3}	59.33	Lutein
5.831	169.38	4.26x10 ⁻³	13.81	Zeaxanthin
6.533	527.83	4.45x10 ⁻³	44.88	Canthaxanthin
7.651	553.82	3.59x10 ⁻³	38.02	Internal Std
14.403				Echinenone
18.453	68.21	7.02x10 ⁻³	9.16	Lycopene
22.278				Neurosporene
31.363	2966.38	3.52x10 ⁻³	199.36	α-carotene
33.870	2854.27	3.86x10 ⁻³	210.64	β-carotene
44.166	524.14	1.59x10 ⁻²	158.86	Phytoene
Totals:			734.05	

Table 10. Carotenoid concentrations of canola seeds from selected T2 6204-SP30021 lines.

			Carote	enoid Conce	entration (µg	g/gFWt.)		
Sample ID	Segreg ratio	Lutein	Canth	Lycopene	a-Carotene	b-Carotene	Phytoene	Total
SP30021	homo	21	ND	ND	ND	ND	ND	21
3390-SP30021-12	homo	44	ND	9	416	578	279	1326
6204-SP30021-1	3:1	41	22	9	223	252	192	744
6204-SP30021-2	15:1	43	24	9	231	283	236	831
6204-SP30021-3	3:1	51	8	6	165	268	65	568
6204-SP30021-5	63:1	56	25	11	292	296	305	992
6204-SP30021-6	no fit	61	47	9	206	218	165	720
6204-SP30021-7	3:1	41	13	8	180	232	160	634
6204-SP30021-8	3:1	41	16	6	68	108	54	299
6204-SP30021-9	>63:1	57	39	10	233	245	245	837
6204-SP30021-10	no fit	33	9	7	165	24	103	343
6204-SP30021-11	3:1	39	7	9	198	266	145	662
6204-SP30021-12	15:1	40	15	10	212	281	172	734
6204-SP30021-13	15:1	52	44	9	207	223	247	788
6204-SP30021-15	no fit	54	20	8	205	291	160	738
6204-SP30021-21	3:1	44	11	8	142	216	126	551
6204-SP30021-24	3:1	47	9	9	149	202	89	509
6204-SP30021-25	15:1	37	33	8	235	257	243	819
6204-SP30021-28	15:1	. 46	11	9	225	288	123	707
6204-SP30021-29	null	18	ND	ND	ND	ND	ND	18
6204-SP30021-30	3:1	33	11	8	203	246	154	659
6204-SP30021-36	15:1	31	17	7	197	275	95	628
6204-SP30021-37	3:1	38	10	7	137	181	117	490
6204-SP30021-41	3:1	42	9	8	250	339	170	821
6204-SP30021-42	3:1	32	4	6	178	216	103	539
6204-SP30021-43	15:1	54	25	10	242	304	151	792
6204-SP30021-44	no fit	48	27	7	226	249	129	692

The initial results demonstrate that as with plants transformed to express crtB alone, plants expressing crtB and crtW contain significant increases in total carotenoid levels. Furthermore, the results show an increase in the levels of canthaxanthin, when compared to the levels obtained from seeds of plants transformed with crtB alone, as

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well as nontransformed control plants. In addition, other products were also produced in plants expressing crtB and crtW. Increased levels of echineone, a reaction intermediate, as well as a putative 4-keto-α-carotene (Figure 17).

M. Carotenoid Analysis of Mature Seeds from crtB Transgenic Arabidopsis Plants

Mature 3390 T2 seeds from Arabidopsis were collected and carotenoid extracts were prepared and analyzed according to the method described in 2I above. These results are shown in Table 11 below. Carotenoid levels are presented as $\mu g/gFW$.

Table 11. Carotenoids of T2 Arabidopsis seeds transformed with crtB.

			Caro	otenoid conc	entration (μg/g I	FWt.)
Sample ID	Lutein	Lycope	ne α-Caro	tene β-Caro	tene Phytoene	Total
— AT001-50 VAR	18	ND	ND	2	ND	20
3390-AT001-1	24	ND	7	20	7	58
3390-AT001-2	57	5	68	139	98	368

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Initial results indicate that seeds from one line of Arabidopsis transformed with napin-crtB had an 18-fold increase in total carotenoid concentration. This line also demonstrate an approximately 70 fold increase in β -carotene levels (Table 11).

N. Carotenoid Analysis of Mature Seeds from crtB Transgenic Corn Plants

Mature 9039 T2 seeds from transgenic corn were collected and carotenoid and tocopherol extracts were prepared and analyzed according to the method described in 2I above. These results are shown in Table 12 below. Carotenoid and tocopherol levels are presented as $\mu g/gFW$.

transformed with crtB. Table 12. Carotenoid and tocopherol concentrations of T2 corn seeds

			Carotenoid o	concentrat	Carotenoid concentration (μg/g FW)		Ţ	copherol c	Tocopherol concentration	n (μg/g FW)	
									1		
Sample ID#	Lutien	Zeaxanthin	Lycopene a-(Carotene	b-Carotene P	Phytoene	Total-C	d-Tocop	g-Tocop	a-Tocop	Total-T
Control-1	7	1.92	QN.	0.44	0.33	4	13	2	63	. 18	82
Control-2	00	2.18	Q	0.43	0.27	∞	18	က	83	19	105
Control-3	∞	2.16	£	0.40	0.41	9	17	ന	83	19	105
Control-4	10	2.33	Ð	0.23	0.32	4	17	n	81	16	101
ZM-S4783	∞	1.70	Ð	0.22	1.25	83	94	5	78	14	96
ZM-S4784	5	1.53	Ð	0.38	0.38	14	22	m	53	9	62
ZM-S4785	5	1.5	Ð	0.13	0.75	36	44	9	113	21	140
ZM-S4789	2	0.79	Q	0.16	0.47	34	37	m	48	7	58
ZM-S4790	'n	0.93	Ð	0.17	0.16	9	10	m	63	11	5 11
ZM-S4791	4	1.42	QN QN	0.32	1.93	54	62	9	103	20	129
ZM-S4795	4	1.05	Q	0.39	0.57	10	17	3	64	13	80
ZM-S4796	æ	0.93	S	0.34	0.48	14	18	ω	72	17	92
ZM-S4801	9	1.95	S	0.29	6.0	16	25	4	88	19	111
ZM-S4805	7	1.65	QZ QZ	09.0	0.39	9	16	-	21	QN ON	23
ZM-S4814	7	1.64	Q	0.12	0.40	5	13	4	78	19	101
ZM-S4815	3	1.24	QN QN	0.27	0.41	7	13	4	91	19	114
ZM-S4816	7	0.92	ON.	0.14	0.62	36	40	3	89	14	98
ZM-S4819	6	3.65	ON ON	0.19	0.81	6	23	9	52	16	75
ZM-S4820	n	0.97	ON	0.11	0.27	4	∞	3	62	15	79
ZM-S4821	6	2.11	OZ.	0.36	0.43	4	16	7	23	18	43
ZM-S4825	c	0.87	QZ	0.20	0.64	29	33	33	65	15	83
ZM-S4826	7	0.72	QX	ON.	0.46	35	38	ന	65	17	85

25558436681 2588888845651 26	83
7	17
4 3 4 5 6 8 4 5 6 8 6 8 6 8 8 8 8 8 8 8 8 8 8 8 8 8 8	63
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0.54 0.96 0.27 0.27 0.47 0.43 0.43 0.43 0.43 0.44 0.34 0.35 0.34 0.34	0.69
0.08 0.19 0.19 0.25 0.26 0.30 0.31 0.31 0.31 0.31 0.31 0.31 0.31	0.72
	2
0.84 1.32 1.07 1.09 0.94 0.95 1.68 1.68 1.68 0.88 1.68 0.88 1.72 2.55 3.98 1.72 2.73 1.81	2.46
w 4 4 4 ≈ 5 ≈ 6 ≈ 6 ≈ 6 ≈ 6 ≈ 6 ≈ 6 ≈ 6 ≈ 6 ≈ 6	∞
ZM-S4827 ZM-S4833 ZM-S4833 ZM-S4833 ZM-S4842 ZM-S4842 ZM-S4849 ZM-S4849 ZM-S4854 ZM-S4855 ZM-S4856 ZM-S4856 ZM-S4865 ZM-S4866 ZM-S4865 ZM-S4868 ZM-S4866 ZM-S4886	ZM-S4889

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0.29 25 3.57 00 00 ZM-S4892 ZM-S4893

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Seeds from 48 transgenic corn lines were analyzed for carotenoid and tocopherol concentrations (Table 12). Total carotenoid concentration was increased up to 5 fold. The major change was phytoene (up to 15 fold increase), with a small increase in b-carotene in some lines. The total carotenoid concentration of transgenic corn seeds was only 5%-10% with respect to that of the *Brassica* seeds transformed with *crtB*. The substantial accumulation of phytoene indicates that phytoene desaturase of corn could be rate limiting. Therefore, the *crtB* effect was basically limited to phytoene biosynthesis and blocked to the later steps of the carotenoid pathway. These wild type corn seeds accumulate some zeaxanthin, but it is only about ¼ that of lutein. In order to produce more zeaxanthin in corn seeds, at least three genes (*crtB*, *crtI*, and *crtZ*)may be required.

O. Expression of Maize phytoene synthase in Arabidopsis

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Mature 9061 T2 seeds from *Arabidopsis* were collected and carotenoid extracts were prepared and analyzed according to the method described in 2I above. These results are shown in Table 13 below and in Figure 18. Carotenoid levels are presented as $\mu g/gFW$.

Table 13: Results of Carotenoid composition determined by HPLC on A.t. 9061 lines

				_		Total-	
Strain ID	Lutein	Lycopene	α-	β-	Phytoene	Major-C	%phytoene
		<u> </u> 	Carotene	Carotene			
AT001	27	0	0	8	0	34	0
9061-	36	4	20	82	15	157	9.7
AT001-1							
9061-	61	6	115	295	115	591	19.3
AT001-2							
9061-	29	4	22	82	24	160	15.0
AT001-3							
9061-	68	5	78	251	60	461	12.9
AT001-4							
9061-	42	6	62	163	63	336	18.7

AT001-5				-			
9061-	46	4	59	167	60	337	17.8
AT001-6							İ
9061-	31	4	17	61	6	119	4.9
AT001-7							
9061-	58	7	80	230	85	460	18.4
AT001-8		<u> </u>	1			}	:
9061-	27	4	24	57	25	138	17.9
AT001-9							
9061-	64	6	82	243	78	473	16.5
AT001-							
10							
9061-	52	6	79	225	72	434	16.4
AT001-							
11						1	
9061-	53	5	74	222	69	424	16.3
AT001-							
12							
9061-	43	5	62	217	55	382	14.3
AT001-				1	,		
13							
9061-	49	5	73	185	83	395	20.9
AT001-							
14							
9061-	26	4	13	60	9	112	8.4
AT001-							
15							
9061-	33	5	52	157	50	297	16.7
AT001-							
16							
9061-	69	6	93	293	86	547	15.7
AT001-							
17							
9061-	41	5	46	164	24	280	8.5
AT001-						1	i
18							_
9061-	55	6	85	206	103	455	22.7
AT001-							
19							
9061-	71	6	116	244	125	562	22.3
AT001-							
20]					
9061-	61	5	88.	226	89	469	18.9

AT001- 21							•
9061- AT001- 22	49	6	66	198	76	395	19.2
9061- AT001- 23	44	4	44	167	22	282	7.8
9061- AT001- 24	59	6	70	219	57	411	13.9
9061- AT001- 25	58	6	65	228	70	426	16.4
9061- AT001- 26	57	6	. 81	212	80	436	18.2
9061- AT001- 27	17	0	8	18	0	. 44	. 0
9061- AT001- 28	55		76	176	65	37.8	17.2

Transgenic Arabidopsis expressing the maize phytoene synthase demonstrate an increase in total carotenoid levels. There are 11 lines of pCGN 9061 that have total carotenoids in excess of 400 ppm. This is higher than the highest pCGN 3390 line which is 381ppm. There are three Arabidopsis lines containing the maize phytoene synthase with total carotenoids between 550ppm and 590ppm.

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Also, transgenic *Arabidopsis* lines containing the maize phytoene synthase produce less phytoene as a percentage of total carotenoids. The pCGN 9061 lines that accumulate the most carotenoids have only 16%-22% phytoene. The pCGN 3390 lines that accumulate the most carotenoids have a percent phytoene range from 25%-31% phytoene. The β-carotene is also significantly increased (Figure 18).

Example 3 Crosses of crtB Plants

A. Transgenic Oil Traits

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To evaluate the high oleic trait of the napin-crtB transgenic plants in conjunction with expression of other oils traits, crosses off 3390-1-6-8 with a mangosteen thioesterase (5266) and a nutmeg thioesterase (3854; see WO 96/23892) were made. Crosses were also made with two low linoleic (LPOO4 and LP30108) varieties. Half-seed analyses of carotenoids and fatty acid composition were conducted on the segregating seeds, and the average of the half seed values are shown below in Tables 13 and 14.

TABLE 13

Carotenoid Levels in Half Seeds Resulting from 3390 Crosses

15	Cross	Lutein	Lycop	ene α-Ca	arotene	B-Carotene	Total
	F1 3390-SP001-1-6-8 x SP30021	21.6	26.2	271.5	413.1	732.4	
	F1 3390-SP001-1-6-8 x 5266-SP3002	21-5-26	18.0	21.7	187.9	284.1	511.7
	F1 3390-SP001-1-6-8 x 5266-SP3002	21-35-2	16.2	22.1	223.0	318.4	579.7
	F1 3390-SP001-1-6-8 x 5266-SP3002	21-35-12	19.5	22.9	196.8	312.8	552.0
20	F1 3390-SP001-1-6-8 x LP30108-19		23.7	22.7	213.4	355.0	614.8
	F1 LP30108-19 x F1 3390-SP001-1-6	6-8	16.4	19.6	156.7	224.5	417.2

TABLE 14

Fatty Acid Composition in Half Seeds Resulting from 3390 Crosses

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<i>L</i>	٠,
\mathbf{r}	_
$\mathbf{\mathbf{\mathcal{U}}}$	~

(3390-SP001-1-6-8 X	0.06	3.84	11.37	62.86	11.06	5.08	3.38
5266-SP30021-35-12)							
(3390-SP001-1-6-8 X	0.06	3.68	11.27	64.80	9.81	5.16	3.04
5266-SP30021-35-2)							
3390-SPOO1-1-6-8 X	0.06	3.66	15.36	60.78	9.30	4.77	3.87
5266-SP30021-5-26							
(3390-SP001-1-6-1 X	2.69	9.80	3.65	64.62	9.72	4.57	1.51
3854-SP30021-20-3)						•	
(3390-SP001-1-6-1 X	6.14	16.35	5.12	54.91	8.23	4.23	2.03
3854-SP30021-20-1)							
(3390-SP001-1-6-1 X	0.07	3.82	11.67	64.52	11.46	3.14	3.08
5266-LP004-2-31)							
(3390-SP001-1-6-8 X	0.05	3.80	1.44	73.66	14.02	3.93	0.67
LP30108-19)							
(LP30108-19 X	0.04	3.31	1.79	79.69	9.26	2.97	0.75
3390-SP001-1-6-8)							
SPOO1-4-10	0.07	4.44	0.99	56.06	21.79	14.31	0.44
3390-SPOO1-1-6-8	0.04	3.46	1.44	77.26	9.30	5.71	0.63
		•					

As the above results demonstrate, a dramatic increase (100 to 200 fold) in α - and β -carotene as well as a 60 fold increase in total carotenoids may be obtained by transformation of plants for expression of an early carotenoid biosynthesis gene under the regulatory control of promoter preferentially expressed in plant seed tissue. This increase in flux primes the pathway for the production of specialty products as described above, and also results in increased production of α -tocopherol (Vitamin E).

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Furthermore, it is evident that the fatty acid composition can also be altered in the transgenic plant seeds. In this manner, seeds can be used to produce novel products, to provide for production of particular carotenoids, to provide high oleic oils, and the like.

All publications and patent applications mentioned in the specification are indicative of the level of skill of those skilled in the art to which this invention pertains.

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All publications and patent applications are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it will be obvious that certain changes and modifications may be practiced within the scope of the appended claims.

IN THE CLAIMS

What is claimed is:

1. A method for altering the carotenoid content in seed of a host corn plant, comprising:

transforming cells of a host corn plant with a construct comprising as operably linked components, a transcriptional initiation region from a gene preferentially expressed in a plant seed, a plastid transit peptide, a DNA sequence derived from a carotenoid biosynthesis gene coding region, and a transcriptional termination region,

producing a transformed host corn plant from said transformed cells, and growing said transformed host corn plant or progeny thereof containing said construct under conditions whereby seed is produced having an altered carotenoid content.

- 2. The method according to Claim 1, wherein said altered carotenoid content is increased.
- 3. The method according to Claim 1, wherein said carotenoid biosynthesis gene is an early carotenoid biosynthesis gene.
- 4. The method according to Claim 3, wherein said early carotenoid biosynthesis gene is selected from the group consisting of geranylgeranyl pyrophosphate synthase, phytoene synthase, phytoene desaturase, and isopentenyl diphosphate isomerase.
- 5. The method according to Claim 1, further comprising introducing into a host corn cell a second construct comprising as operably linked components, a promoter functional in a corn seed cell, a second carotenoid biosynthesis gene, and a transcriptional termination region functional in a corn seed cell.

- 6. The method according to Claim 5, wherein said second carotenoid biosynthesis gene is selected from geranylgeranyl pyrophosphate synthase, phytoene synthase, phytoene desaturase, isopentenyl diphosphate isomerase, β -carotene hydroxylase, the astaxanthin biosynthesis enzyme encoded by crtW, and ϵ -hydroxylase, or wherein said secondary gene results in inhibition of transcription of an endogenous plant gene encoding lycopene ϵ -cyclase, lycopene β -cyclase or phytoene desaturase.
- 7. A method for the alteration of carotenoid content in the endosperm of a host plant, comprising:

transforming cells of a host plant with a construct comprising as operably linked components, a transcriptional initiation region from a gene preferentially expressed in a plant seed endosperm, a plastid transit peptide, a DNA sequence derived from a carotenoid biosynthesis gene coding region, and a transcriptional termination region,

producing a transformed host corn plant from said transformed cells, and growing said transformed host corn plant or progeny thereof containing said construct under conditions whereby seed is produced having an endosperm having an altered carotenoid content.

- 8. The method according to Claim 7, wherein said carotenoid content is increased.
- 9. The method according to Claim 7, wherein said carotenoid biosynthesis gene is an early carotenoid biosynthesis gene.
- 10. The method according to Claim 7, wherein said early carotenoid biosynthesis gene is selected from the group consisting of geranylgeranyl pyrophosphate synthase, phytoene synthase, phytoene desaturase, and isopentenyl diphosphate isomerase.

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- 11. The method according to Claim 7, further comprising introducing into a host corn cell a second construct comprising as operably linked components, a promoter functional in a corn seed cell, a second carotenoid biosynthesis gene, and a transcriptional termination region functional in a corn seed cell.
- 12. The method according to Claim 11, wherein said second carotenoid biosynthesis gene is selected from geranylgeranyl pyrophosphate synthase, phytoene synthase, phytoene desaturase, isopentenyl diphosphate isomerase, β -carotene hydroxylase, the astaxanthin biosynthesis enzyme encoded by crtW, and ε -hydroxylase, or wherein said secondary gene results in inhibition of transcription of an endogenous plant gene encoding lycopene ε -cyclase, lycopene β -cyclase or phytoene desaturase.
- 13. A method for screening transformed corn seeds or transformed endosperms, comprising:

transforming a host corn plant with an expression cassette comprising as operably linked components, a transcriptional initiation region from a gene preferentially expressed in a plant seed, a transit peptide, a DNA coding sequence of at least one carotenoid biosynthesis gene, and a transcriptional termination region, and

selecting said transformed seeds or transformed endosperms exhibiting a yellow, orange or red color.

- 14. The method according to Claim 13, wherein said early carotenoid biosynthesis gene encodes an enzyme selected from the group consisting of geranylgeranyl pyrophosphate synthase, phytoene synthase, phytoene desaturase, and isopentenyl diphosphate isomerase.
- 15. The method according to Claim 13, further comprising introducing into a host corn cell a second construct comprising as operably linked components, a promoter

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functional in a corn seed cell, a second carotenoid biosynthesis gene, and a transcriptional termination region functional in a corn seed cell.

- 16. The method according to Claim 15, wherein said second carotenoid biosynthesis gene is selected from geranylgeranyl pyrophosphate synthase, phytoene synthase, phytoene desaturase, isopentenyl diphosphate isomerase, β -carotene hydroxylase, the astaxanthin biosynthesis enzyme encoded by crtW, and ϵ -hydroxylase, or wherein said secondary gene results in inhibition of transcription of an endogenous plant gene encoding lycopene ϵ -cyclase, lycopene β -cyclase or phytoene desaturase.
 - 17. A nucleic acid sequence encoding a phytoene synthase from corn.
- 18. The nucleic acid sequence according to Claim 17, wherein said sequence is the sequence set forth in Figure 19.
- 19. A method for altering carotenoid composition in a seed from a host plant, said method comprising transforming said host plant with a construct comprising as operably linked components, a transcriptional initiation region from a gene preferentially expressed in a plant seed, a plastid transit peptide, a nucleic acid sequence encoding a carotenoid biosynthesis gene from eukaryotic source, and a transcriptional termination region.
- 20. The method according to Claim 19, wherein said carotenoid biosynthesis gene is phytoene synthase.
- 21. The method according to Claim 19, wherein said carotenoid biosynthesis gene is from corn.

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22. The method according to Claim 19, wherein said carotenoid biosynthesis gene is encoded by the nucleic acid sequence set forth in Figure 19.

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3

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Met Thr Gly Phe Pro Val Lys Lys Val Asn Thr Asp Ile Thr Ser Ile 35 40 45

Thr Ser Asn Gly Gly Arg Val Lys Cys Met Thr Asn Phe Leu Ile Val 50 55 60

Val Ala Thr Val Leu Val Met Glu Leu Thr Ala Tyr Ser Val His Arg
65 70 75 80

Trp Ile Met His Gly Pro Leu Gly Trp Gly Trp His Lys Ser His His 90 95

Glu Glu His Asp His Ala Leu Glu Lys Asn Asp Leu Tyr Gly Leu Val 100 105 110

Phe Ala Val Ile Ala Thr Val Leu Phe Thr Val Gly Trp Ile Trp Ala 115 120 125

Pro Val Leu Trp Trp Ile Ala Leu Gly Met Thr Val Tyr Gly Leu Ile 130 135 140

Tyr Phe Val Leu His Asp Gly Leu Val His Gln Arg Trp Pro Phe Arg 145 150 155 160

Tyr Ile Pro Arg Lys Gly Tyr Ala Arg Arg Leu Tyr Gln Ala His Arg

165 170 175

Leu His His Ala Val Glu Gly Arg Asp His Cys Val Ser Phe Gly Phe 180 185 190

Ile Tyr Ala Pro Pro Val Asp Lys Leu Lys Gln Asp Leu Lys Met Ser 195 200 205

Gly Val Leu Arg Ala Glu Ala Gln Glu Arg Thr 210 215

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Met Thr Gly Phe Pro Val Lys Lys Val Asn Thr Asp Ile Thr Ser Ile 35 40 45

Thr Ser Asn Gly Gly Arg Val Lys Cys Met Ser Ala His Ala Leu Pro 50 60

Lys Ala Asp Leu Thr Ala Thr Ser Leu Ile Val Ser Gly Gly Ile Ile 65 70 75 80

Ala Ala Trp Leu Ala Leu His Val His Ala Leu Trp Phe Leu Asp Ala 85 90 95

Ala Ala His Pro Ile Leu Ala Ile Ala Asn Phe Leu Gly Leu Asn Trp 100 105 110

Leu Ser Val Gly Leu Phe Ile Ile Ala His Asp Ala Met His Gly Ser 115 120 125

Val Val Pro Gly Arg Pro Arg Ala Asn Ala Ala Met Gly Gln Leu Val 130 135 140 ·

Leu Trp Leu Tyr Ala Gly Phe Ser Trp Arg Lys Met Ile Val Lys His 145 150 155 160

Met Ala His His Arg His Ala Gly Thr Asp Asp Pro Asp Phe Asp 165 170 175

His Gly Gly Pro Val Arg Trp Tyr Ala Arg Phe Ile Gly Thr Tyr Phe 180 185 190

Gly Trp Arg Glu Gly Leu Leu Pro Val Ile Val Thr Val Tyr Ala 195 200 205

Leu Ile Leu Gly Asp Arg Trp Met Tyr Val Val Phe Trp Pro Leu Pro 210 220

Ser Ile Leu Ala Ser Ile Gln Leu Phe Val Phe Gly Thr Trp Leu Pro 225 230 235 240

His Arg Pro Gly His Asp Ala Phe Pro Asp Arg His Asn Ala Arg Ser . 250 255

Ser Arg Ile Ser Asp Pro Val Ser Leu Leu Thr Cys Phe His Phe Gly 260 265 270

Gly Tyr His His Glu His His Leu His Pro Thr Val Pro Trp Trp Arg 275 280 285

Leu Pro Ser Thr Arg Thr Lys Gly Asp Thr Ala

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Thr Lys Arg Pro Ala Ala Arg Arg Trp Met Pro Cys Ser Leu Leu Gly
35 40 45

Leu His Pro Trp Glu Ala Gly Arg Pro Ser Pro Ala Val Tyr Ser Ser 50

Leu Ala Val Asn Pro Ala Gly Glu Ala Val Val Ser Ser Glu Gln Lys
65 75 80

Val Tyr Asp Val Val Leu Lys Gln Ala Ala Leu Leu Lys Arg Gln Leu 85 90 95

Arg Thr Pro Val Leu Asp Ala Arg Pro Gln Asp Met Asp Met Pro Arg 100 105 110

Asn Gly Leu Lys Glu Ala Tyr Asp Arg Cys Gly Glu Ile Cys Glu Glu 115 120 125

Tyr Ala Lys Thr Phe Tyr Leu Gly Thr Met Leu Met Thr Glu Glu Arg 130 135 140

Arg Arg Ala Ile Trp Ala Ile Tyr Val Trp Cys Arg Arg Thr Asp Glu 145 150 155 160

Leu Val Asp Gly Pro Asn Ala Asn Tyr Ile Thr Pro Thr Ala Leu Asp 165 170 175

Arg Trp Glu Lys Arg Leu Glu Asp Leu Phe Thr Gly Arg Pro Tyr Asp 180 185 190

Met Leu Asp Ala Ala Leu Ser Asp Thr Ile Ser Arg Phe Pro Ile Asp 195 - 200 205

Ile Gln Pro Phe Arg Asp Met Ile Glu Gly Met Arg Ser Asp Leu Arg 210 215 220

Lys Thr Arg Tyr Asn Asn Phe Asp Glu Leu Tyr Met Tyr Cys Tyr Tyr 225 230 235 240

Val Ala Gly Thr Val Gly Leu Met Ser Val Pro Val Met Gly Ile Ala 245 250 255

Thr	Glu	Ser	Lys 260	Ala	Thr	Thr	Glu	Ser 265	Val	Tyr	Ser	Ala	A1a 270	Leu	Ala
Leu	Gly	Ile 275	Ala	Asn	Gln	Leu	Thr 280	Asn	Ile	Leu	Arg	Asp 285	Val	Gly	Glu
Asp	Ala 290	Arg	Arg	Gly		Ile 295	Tyr	Leu	Pro	Gln	Asp 300	Glu	Leu	Ala	Gln
Ala 305	Gly	Leu	Ser	Asp	Glu 310	Asp	Ile	Phe	Lys	Gly 315	Val	Val	Thr	Asn	Arg 320
Trp	Arg	Asn	Phe	Met 325	Lys	Arg	Gln	Ile	Lys 330	Arg	Ala	Arg	Met	Phe 335	Phe
Glu	Glu	Aļa	Glu 340	Arg	Gly	Val	Asn	Glu 345	Leu	Ser	Gln	Ala	Ser 350	Arg	Trp
Pro	Val	Trp 355	Ala	Ser	Leu	Leu	Leu 360	Tyr	Arg	Gln	Ile	Leu 365	Asp	Glu	Ile
Glu	Ala 370	Asn	Asp	Tyr	Asn	Asn 375	Phe	Thr	Lys	Arg	Ala 380	Tyr	Val	Gly	Lys
Gly 385	Lys	Lys	Leu	Leu	Ala 390	Leu	Pro	Val	Ala	Tyr 395	Gly	Lys	Ser	Leu	Leu 400
Leu	Pro	Суѕ	Ser	Leu 405	Arg	Asn	Gly	Gln	Thr 410	Xaa	Pro	Pro	Glu	Lys 415	Leu
Gln	Asp	Pro	Pro 420	Arg	Asp	Xaa	Arg	Ala 425				•		•	

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780	GTTATCTGCC	GCGGGCCGCT	CGATGCGCAT	ATATTGTGGA	ATTGCTCGCG	GTTGACCAAT ATTGCTCGCG ATATTGTGGA CGATGCGCAT GCGGGCCGCT	
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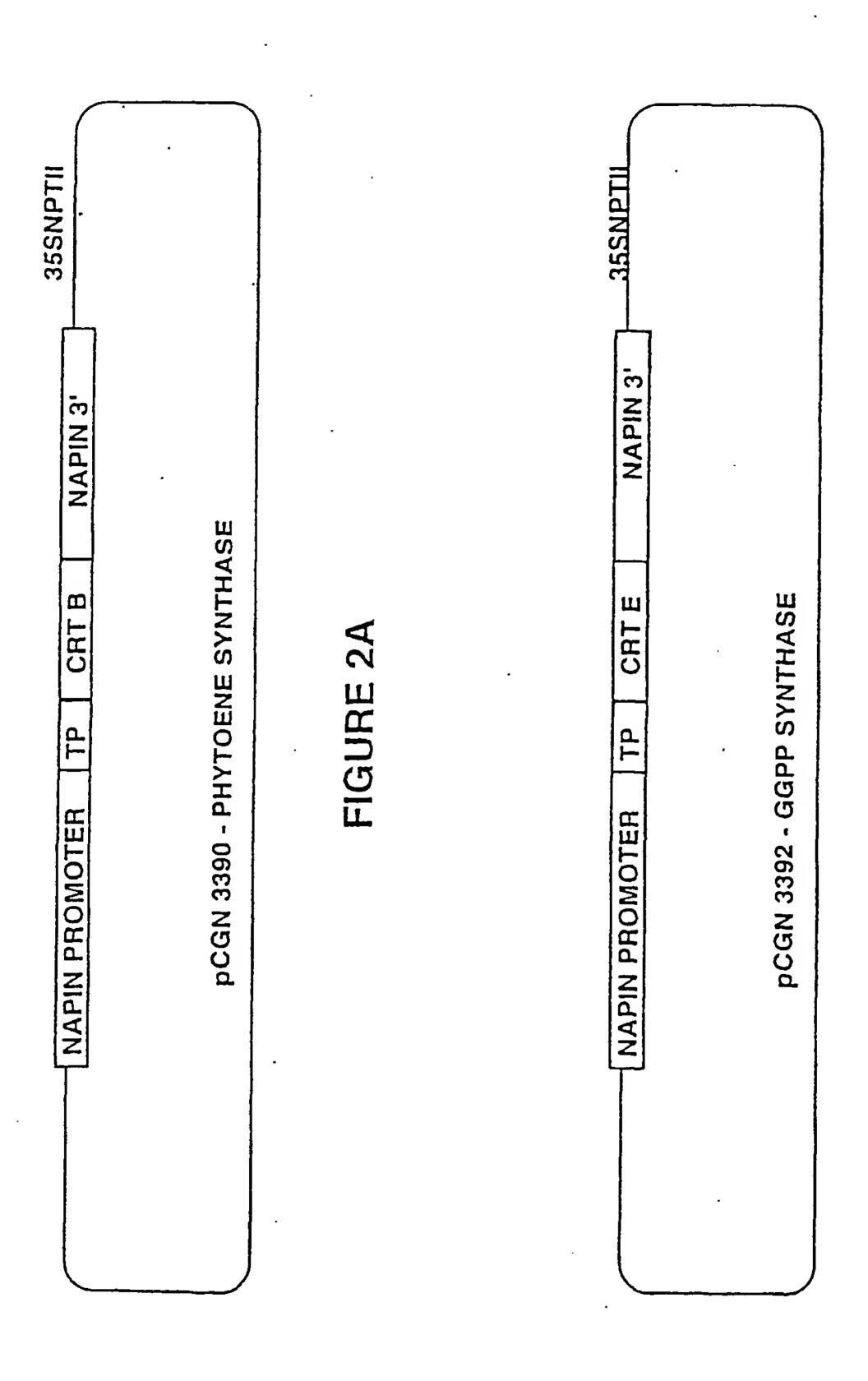


FIGURE 2B

35SNPTII PCGN 9010 - PHYTOENE DESATURASE NAPIN PROMOTER

FIGURE 2C

35SNPTII PCGN 9009 - PHYTOENE SYNTHASE + PHYTOENE DESATURASE NAPIN PROMOTER NAPIN 3' CRT B NAPIN PROMOTER

FIGURE 2D

35SNPTII

35SNPTII NAPIN PROMOTER CRT B | NAPIN 3' TP NAPIN PROMOTER

PCGN 9002 - PHYTOENE SYNTHASE + ANTISENSE EPSILON CYCLASE

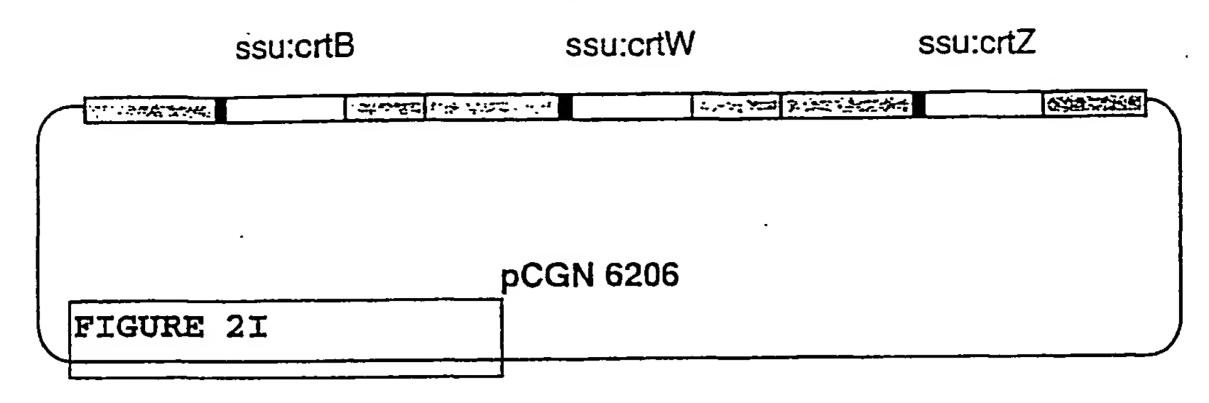
FIGURE 2E

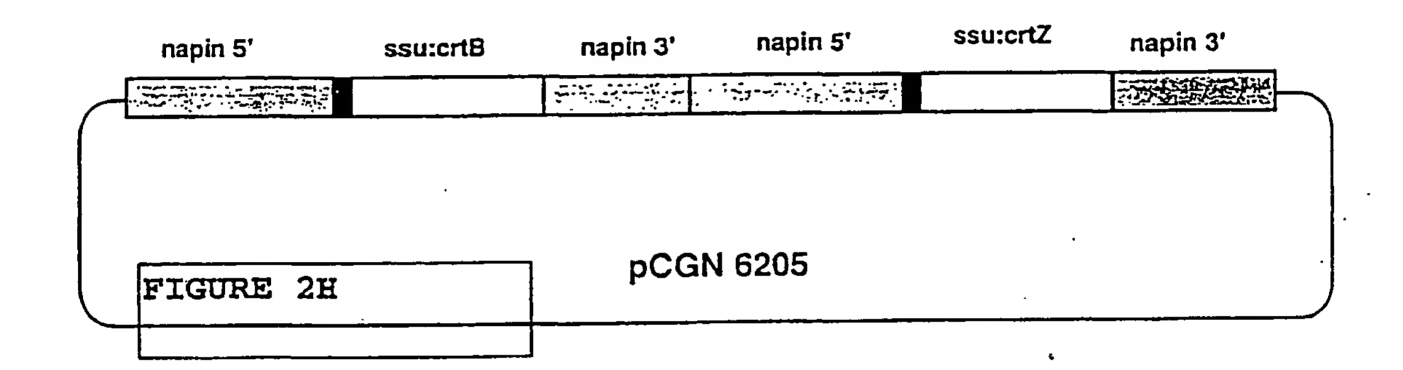
CRT B | NAPIN 3' NAPIN PROMOTER

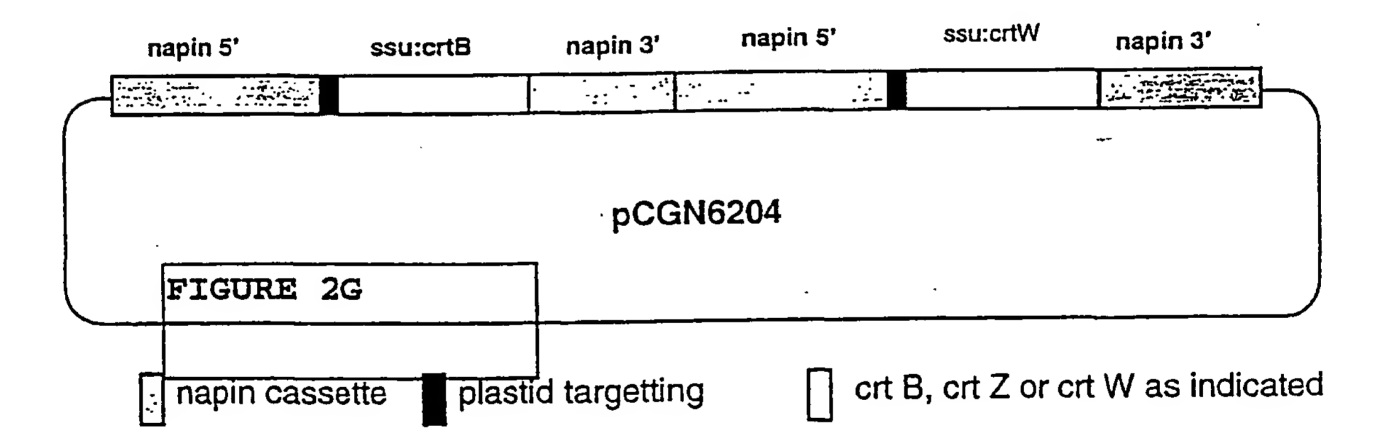
NAPIN **B-CYCLASE** NAPIN PROMOTER

+ ANTISENSE BETA CYCLASE **PCGN 9017 - PHYTOENE SYNTHASE**

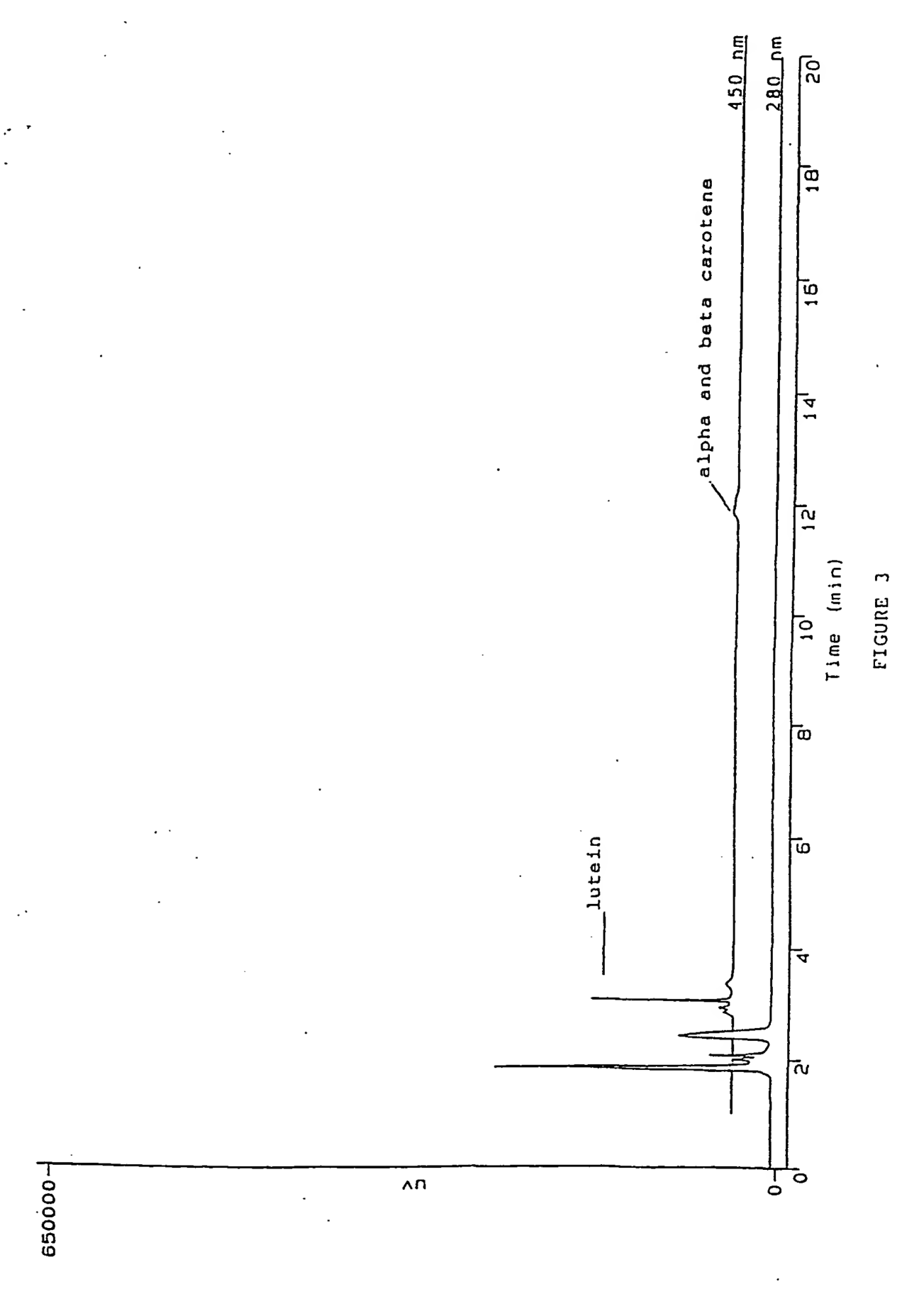
FIGURE 2F











20

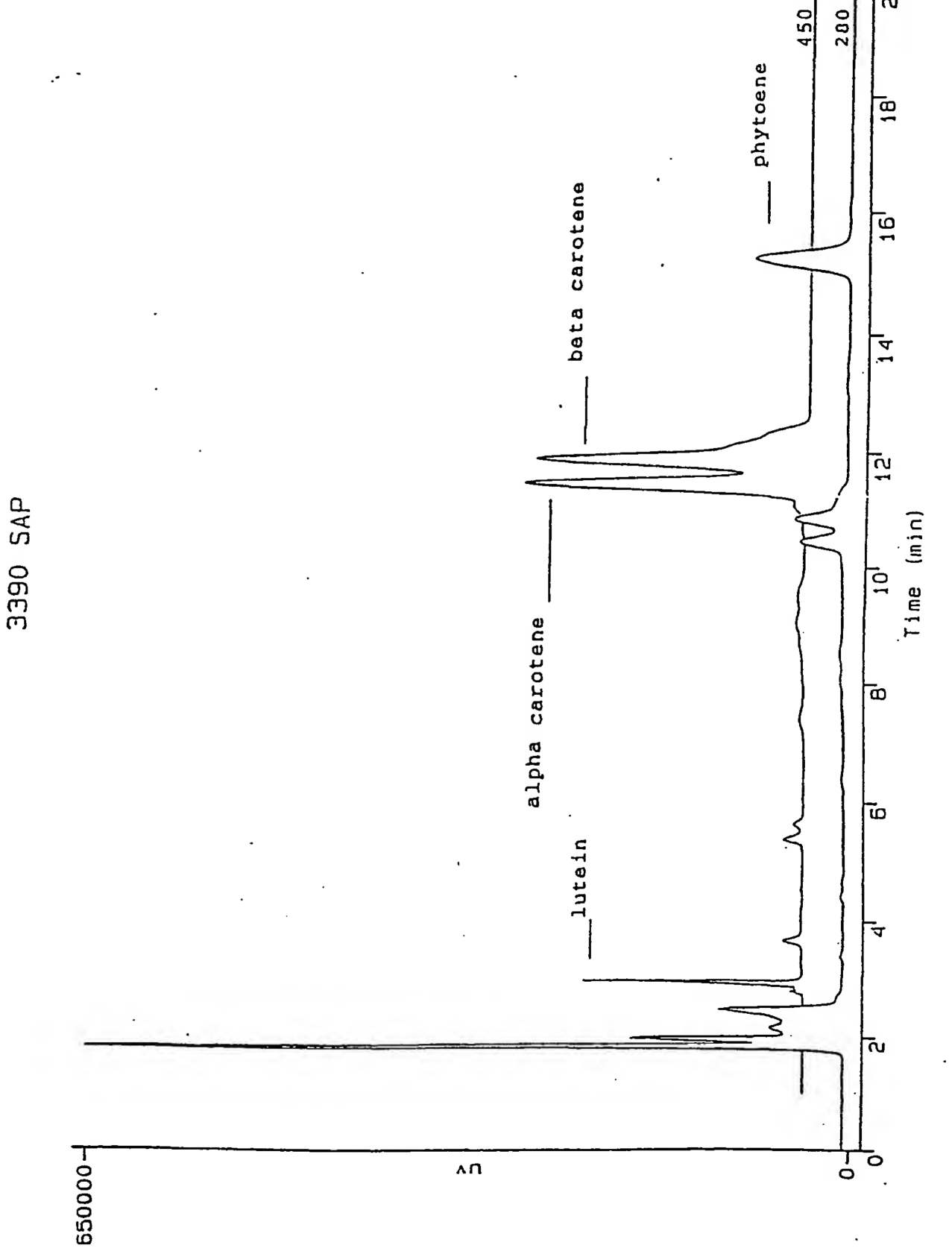


FIGURE 4

18:1 vs 18:2 and 18:3 in 3390s

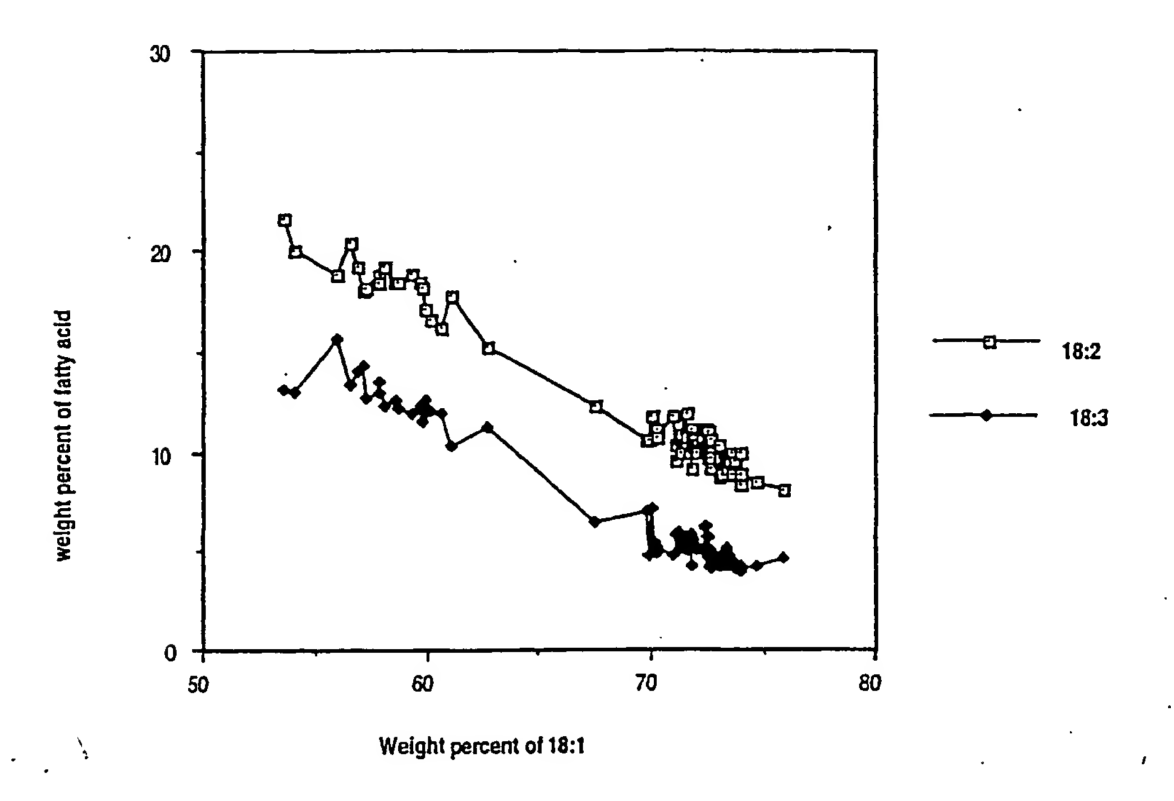


FIGURE 5

18:1 vs 16:0, 18:0 & 20:0 in 3390s

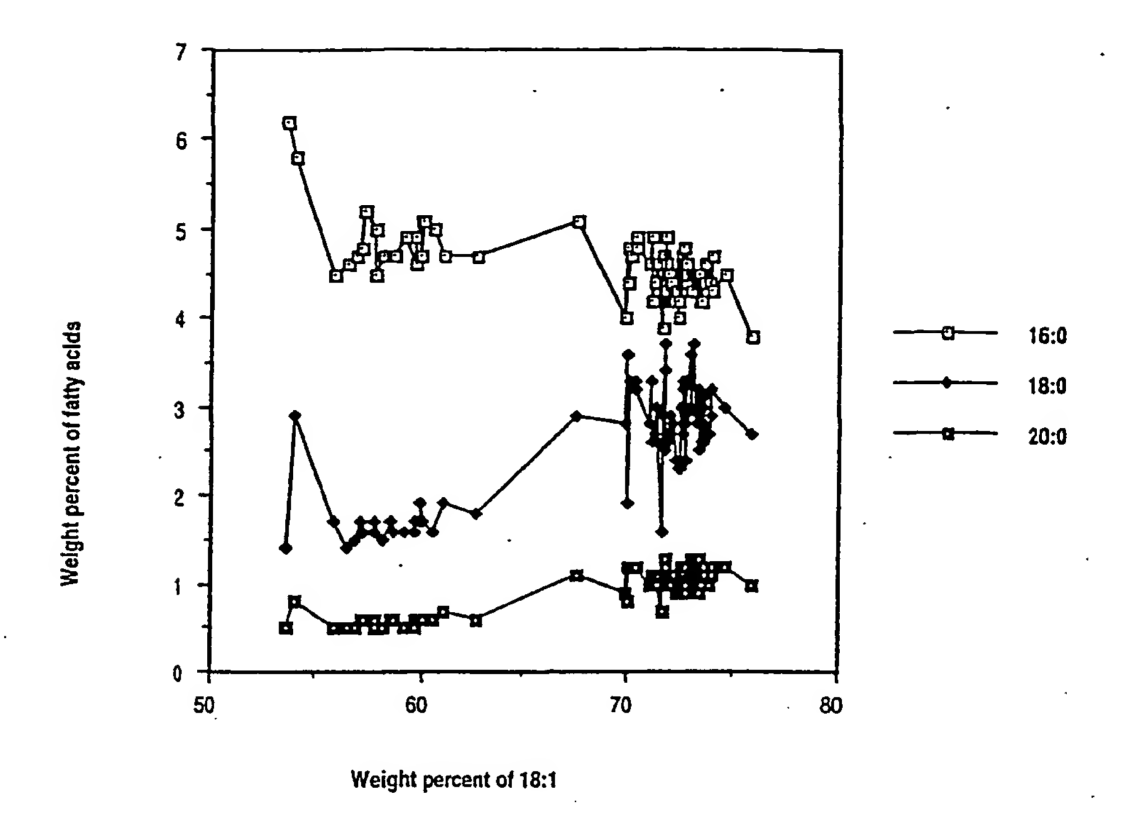


FIGURE 6

18:0 vs 20:0 in 3390s

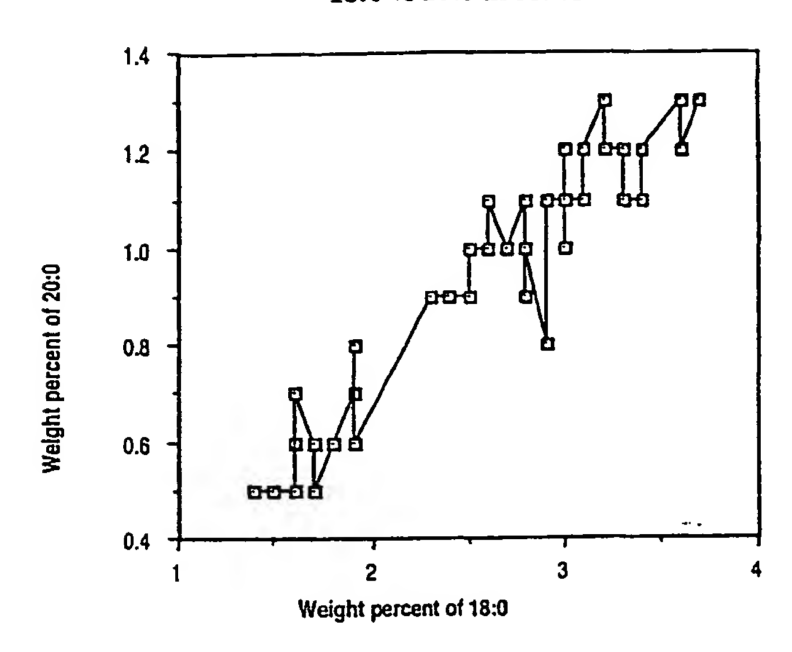
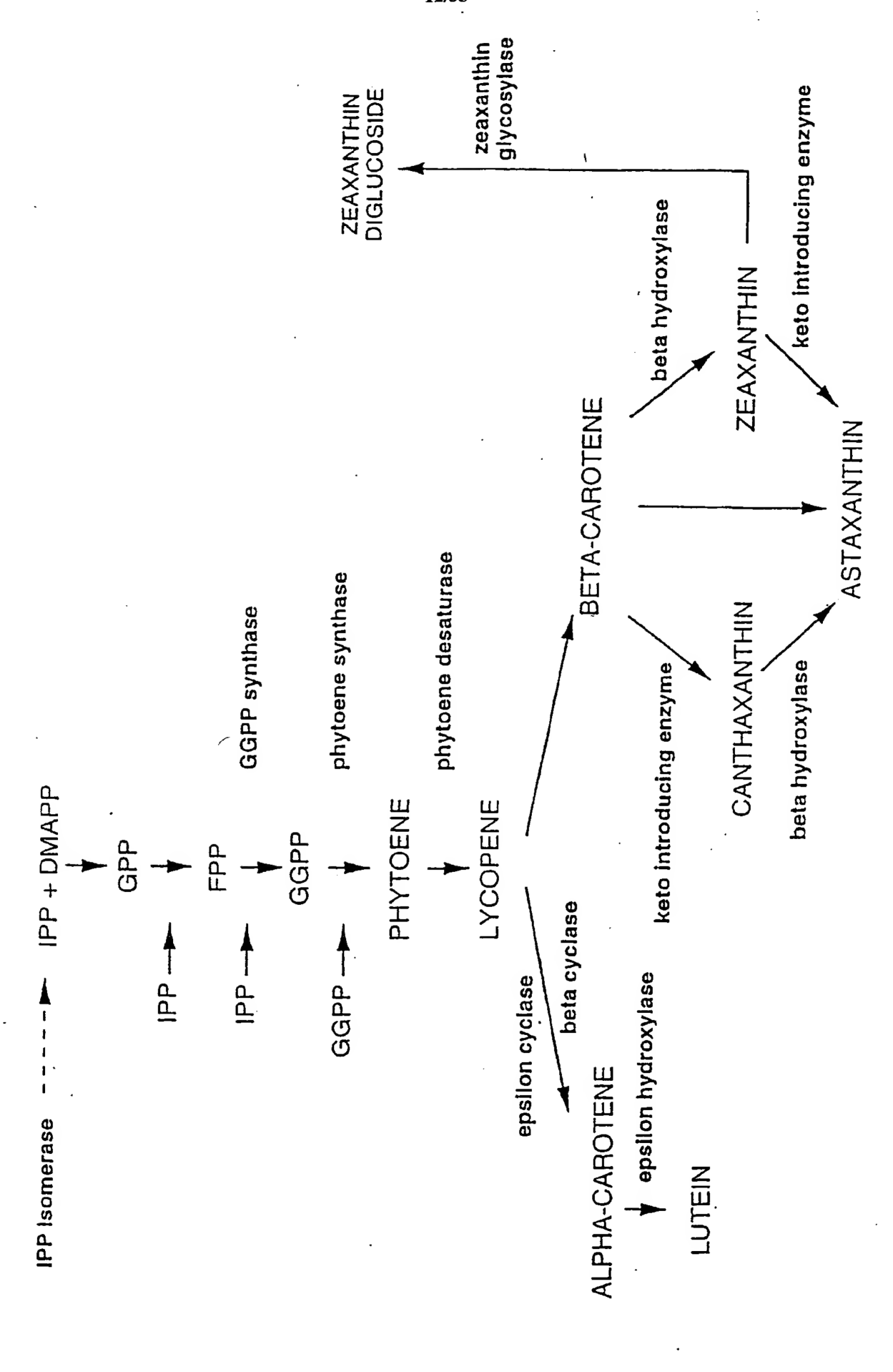


FIGURE 7



FIGURE

 $\boldsymbol{\omega}$

```
PCT/US01/15264
 WO 01/88169
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                                     >XbaI
                                                 >XhoI
                                                >SphI
            >HinfI
                               >HaeIII
                   20
                       >EcoRI
                      >PstI
                >EcoRV
        >2stXI
                                                         120
                                     100
                   80
  >EinfI
                                              >Sau3AI
                                   >HinfI
       >PstI
                                     1160
                                                         180
                  140
                                   >Sau3AI
                  200
                                                         240
                                                         300
                  260
                                     280
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                                           >SacI
                     >HincII
                                         >AluI
                    >SalI
                                                         360
                  320
  GTCGTGCTTATGGAAGAGTTAGTCGACGTTTACTTCACGAGGAGCTCTTGAGGAGGTGTG
                                                >AluI
  >HinfI
                       >AluI
                                                         420
                                      400
                  380
  TGGAGTCAGGTGTCTCGTATCTTAGCTCCAAAGTTGAGAGCATAACAGAAGCTCCTGATG
>HaeIII
                                      460
                                                         480
                  440
                                                 >HinfI
```

FIGURE 9

WO 01/88169

GA

PCT/US01/15264

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>EcoRI	>Pati	V 90	210 TTGGGAAGAT	>Hincii 	>Sali 330 1 TAGTCGACOT	450 TGAACAAAAG	ggAGGTT	כאדפככד
		>ECOR 80 * * TGGATAT	200 200 ACTACGGTGT 1		320 , ATGGAAGAGT	440	560 •	680 TTCTCTACGC
		>Bs 70 *	190		310 1 orcorderr	• 11aaIII 0 430 • 430	550 • atccaaactg	670 TATCCAACOT
	· Ior	> Ha	180		310 1 TATTACCATT GGTCGTGCTT	AGCTCCTGAT G	_	Hapi Alui 650 670 100 600 670 TCCGGAGCTT AGAAGCTGAA TATCCAACGT
	xhor.	Sai Sphi	170		290 ATGACAATCC TA	410 410		Hapi 650
		>Haelli >Xbal			.280 * TATCTGGACG ATC	400	·	640 AACGAGAAAA
		30 30 CTATAGGGCG A		GTTACGACTT A	270 1000			630 • AGATTATACA
		>Hinfl 20 ATACGACTCA CTA	Aluī 140	AATCAGCTAA GT	260 270 TTTGGAGAGA TACCCTTGTG	380	STGTCTCGTA TCTTRUCTCC >AluI >AluI 500 510 CAGCTTCTGG GAAGCTCTTG	620 TGGATTACAG A
		10 aranattata Atac	>BglI >HinfI 130	TTGGCGGCTG AAT	250 * ATTGAGCATG TT		Grogagread Greferera Terraderies Alui Al	610 ATGGTGTTCA TO

FIGURE 10

											10,	33			
	840	TATATCCCA			960	rccalantac		•	1080	ла соасла				1200	OTTATCO
	830	AAATGGTC 7		>Aluī	- 6 - 6 - 6	arcravage '			1070	CAGAAAG GA				1190	CTGTGTGAAA TTGTTATCCG
••	820	TTACOANG AC	>Bglif	SaulaI ,	940	ם מממארכדד דע		>HaeIII	1060	TINICOC CAC			I	1180	crorrre cre
	810	TACTAAAG AC	٨	3	930	ATTCAGTE GT			1050	AGCTTGGGAT ACTTTATGGC CACCAGAAAG GAAACGACAG			*AluI	1170	ATCATEGICA TACCIOTITC
.	800	CTTGTTGAGA TTAGAGACAC TCGGAATCCG AATACTAAAG ACTTACGAAG AGGAATGGTC TTATATCCCA	•		920	Crochachgo craficaote Groadatere torchonde tecanatae	*A1uI	- Hindiii	1040					1169	derreacorn Arc.
	790	FAGAGACAC TO			910				1030	ACCANTA TIT	>Alui	IIIpujik :	*Kpn I	1150	TCGGTACCAA GCT
	780	TTGTTOAGA T	>Aluī	→ ₽8 ¢.	006	TOCAGCTAGC ATGOTTCATC			1020	SACACATC AAC		11 > Sac I	I *Aluf	1140	
-	770				890				. 1010	cerrene en		SpeI > Bamil	>5Au3AI	1130	ידאכדאקד מם
	760	GATTTGCTTA AAAAGAAGCT		>Hinfi	880	, hanatetea e			1000	ACCACTA CTT		*	: >!{aefii	1120	- -TGGCGGC CG
	750		• •	^	870	ACGGAACAA A			066	AAAACAT GAG			Y BetXI	1110	TCCAGCA CAC
	740	MAAGATGT CA			960	CTTOCCAAA C			086	CTAATAT ACT			> EcoRI	1100	TAAGCCG AAT
	730	TOTOTTOCTT CANAGATGT CATGCCCTTT			850	OTABOTOTT CETTOCCANA CACGGAACAA AABAATCTCO CETTTOOTOC			910	GCATCAGTCA TCOCTAATAT ACTAAAACAT GAGACCACTA CTTCCTTCAC CAGACACATC AACACCAATA TTTCAAGACA				0601	AND THE THETANGERS ANTHERAGES CACTEGERS COTTACTAGT GOALECGAGE

GGCTGCCTAA

CTCACAATTC CACACAT ACGAGCCGGA AGCATAAAGT GTAAAGCCTG

1240

1230

1220

1210

>Msp.

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FIGURE

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.420 240 480 180 300 360 CCTCAGCCGA GAGCTCATCT TGCAACTATC CATCCCCGCC ATCTTGTACC 120 CAACCTCGAG ATCAAAGAAC GAAGAACTCC GAGAAACAGC AAAATATCCA TCCCAAAACA CTATAGCGTT AAGGATGAAC TTCGGCTTCT GTTCCCTTCG TCATGATCTC CCAGGTCCGG CAAAGAACGC TAACCAGTCG TCTCTCCACA GGTTCCAGGT AGGTACCGCA TGTGCTGGAA CCGAAGAACA CTCTGTCTCC GCCTCTCAAT AGGCCACAAG CCTCTCAAGC CGTTGTTGTT ACCACCAC CTCCTAGTAG CATCCAAATC AAGCTTCAGC GGAGCAGCTG CAAGAGTCCT AGCAACCATG GTATTAGAAG CATGCGAGAA GAGCGACAGC CTCGACGACA AGAAACCATG CCAGTAACGC AAATTGTTGA TCATCTTAGC AAGAGGAACA GAGCTCGGAT CCACTAGTAA CGGCCGCCAG

840 * GTTAACCTCC .096 ***** CATGAACACC AGTAGCGTCG 720 900 099 AGACCCGGCC TAGCCACAAG AGACGTCTCT 780 TGATACCATA AGCTACTTGA 540 CATAGGGATC CCAAATGTTT CAGCCTCGCA GGAACGTAGG GATCTTGCTG CAACAGTGGA AAGACCGGTA AAGGACCTCC CTCTCCAGTC TTGAAAAACC TCACTGCAGA ATCGCGTACA ACGTTGATCC AGATGCTTAT ACCTCAGCAA ACCAAGCATC TACCCAGGGT TATAAGGCTT ATCATACTGA GTACCGCCAA TGCCAACAC TCTTTGAGGC ACACAACGCT CGTCTTCCTC AATCCGCTTC ACCATCCTCT CTTGGATATC TTCCATCTTC TICCGCTCTT TAACCICAGG GTACGCGTCA ATCTTATCCA CATCAAACGG GTGACCATCG AGAACAACAG AAGCCTGAAT CTTCACACCG TCAAGAAAGA TCCTGTTGGA AGAAACGGC

1440

CTTCAAACCA

TYGCTCTTTT

CCGATTCTGA

AGAGCACTAC TAGCACTAGT GATACTACTA

1020

CTGAGCCACG 1320 1380 TCCTTCTTGG TTTCAGGAAC AAGATCCAAA 1080 1260 CAMCTTTGTT CAGGGAAGTG GAGGCAGTCT 1200 CAAACTCCGT AGTTGTTAGG CCAAATGAGT 1140 GATGCACTTC GGACAGGTCC CGGAGACTTG AGGTGGTGTC CACCGTTGGT CGTAAGGCCG AGTCCAGCTT GCGCCGGACC TGGAATCTAA TTCACTCTCC GCTAGATCAA TCGTTTAAAG GAAGGTCCAA GTCGAGATTC TTGGGGGAAG GATCGATGGA GCAGACAGAG GCTAAACCAG CGGGGCCGCC GCCAACGATA AGCAAGTCCA TGGCTTCGAA CTCGTCAACC TCGTGAACCA CGTCAGTGAC TTTAGCCTGA TGAAGCATCT TGGATTTGAG CTGTTTACGG TTTTCGGAGC CGTCGTTGAT GTAGACGACG

1560 AGGGATGAAA CATGAAACTG

AGCTTAACCC TTGAAGGATT TGGACTTAAT

GTATCCATCG AACTCGAGCT TGTTGGGTGT TTTCAACAGA

CACTGGCGGC CGCTCGAGCA TGCATCTAGA

	Total	26.3	38.9	654.3	1150.4	983.5	805.3	1077.3*	1298.1	4	1425.9	1423.7	87.0	1186.0*	1197.0*	384.1	9.686	28.2	872.5	1001.3	1266.0	1029.1	888.3	582.4	48.0	1134.4*	853.7	934.1	939.2*	1048.5*
n (µg/gFW)	B-Carotene	1.9	4.9	385.7	721.4	580.9	463.3	659.4	. 797.0	941.4	904.3	949.2	22.8	714.5	738.8	216.0	611.6	1.2	527.4	614.3	759.3	613.2	544.9	334.2	4.9	663.9	520.4	580.9	584.4	686.3
concentration (µg/gFW)	α-Carotene	ND	ΩN	229.0	372.4	352.9	306.1	370.6	445.1	494.9	468.4	394.2	12.6	409.8	400.0	98.1	320.0	NO	283.1	324.9	449.0	346.1	285.0	187.9	ND	358.6	272.1	309.1	311.2	299.8
Carotenoid	Lycopene	ND	ON ON	6.1	6.2	3.9	4.9	10.5	9.1	7.4	11.3	11.9	ND	9.5	10.2	3.9	8.9	ND	6.4	9.1	8.1	7.6	6.3	4.1	N N	10.9	7.3	12.2	.9.3	9.8
	Lutein	24.4	34.0	33.5	50.4	45.8	31.0	36.8	ø.	51.2	41.9	68.4	51.6	52.2	48.0	66.1	49.1	27.0	55.6	53.0	49.6	62.2	52.1	56.2	43.1	71.0	53.9	31.9	34.3	52.6
Segregation	ratio			3:1	15:1	no fit	3:1	3:1	15:1	15:1	no fit	>63:1	Innll	3:1	3:1	3:1	3:1	llau	. 3:1	3:1	>63:1	3:1	3:1	3:1	lluu	3:1	no fit	3:1	3:1	3:1
	Sample ID #	SP30021 control 1	SP30021 control 2	T2 3390-SP30021-1	T2 3390-SP30021-2	T2 3390-SP30021-3	T2 3390-SP30021-4	T2 3390-SP30021-5	T2 3390-SP30021-6	T2 3390-SP30021-7	T2 3390-SP30021-8	T2 3390-SP30021-9	T2 3390-SP30021-10	T2 3390-SP30021-11	3390-S	T2 3390-SP30021-13	T2 3390-SP30021-14	T2 3390-SP30021-15	T2 3390-SP30021-16	T2 3390-SP30021-17	T2 3390-SP30021-18	T2 3390-SP30021-19	T2 3390-SP30021-20	T2 3390-SP30021-21	T2 3390-SP30021-22	T2 3390-SP30021-23	T2 3390-SP30021-24	3390-SP30021-	T2 3390-SP30021-26*	T2 3390-SP30021-27

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1432.4	1375.9	1024.8	912.9	140.5	875.2	1409.6	732.9	1434.9	977.3*	963.7	40.9	796.0	1189.3*	630.9	437.9	602.5	965.8	675.0	1053.1	1162.7
7.706	822.9	598.4	527.1	55.0	546.9	808.3	461.0	829.9	625.9	576.1	3.6	505.2	719.3	352.9	219.8	368.3	570.5	401.5	677.2	669.3
446.3	459.4	356.9	302.5	31.4	283.1	502.3	224.5	538.5	291.1	309.0	S	251.1	414.1	230.3	128.4	211.2	312.5	225.2	346.0	439.5
10.0	8.5	5.8	7.3	2.3	8.9	12.1	8.1	11.0	10.0	8.1	QX	1.8	8.4	5.1	5.6	1.4	3.7	3.0	1.6	1.5
68.4	85.1	63.7	76.0	51.8	•	86.9	39.3	•	50.3	•	37.3		47.5	42.6	83.3	21.6	79.1	45.3	28.3	52.4
no fit	>63:1	3:1	3:1	null	3:1	>63:1	3:1	15:1	3:1	3:1	null .	3:1	3:1	3:1	no fit	3:1	3:1	3:1	15:1	Ното
-c	3390-SP30021	33	3390-SP30021	3390-SP30021-	33	3390-SP30021-	3390-SP30021-	3390-SP30021-	3390-SP30021-3	3390-SP30021-	3390-SP30021-3	33	33	3	3	3	3390-SP30021-4	3	ä	ξ.

arotenoid concentration (µg/gFW)

	Cogregation		arotenoid c	Carotenoid concentration (µg/grw)	(µg/grw)	
Sample ID #	status	Lutein	Lycopene	α-Carotene	β-Carotene	Total
T3 3390-SP001-4-12	Homo	43.9	17.2	282.1	636.8	980.0
T3 3390-SP001-5-7	Het	50.7	6.3	190.6	386.8	634.4
T3 3390-SP001-5-12	Homo	45.5	19.5	255.9	633.4	954.3
T3 3390-SP001-11-6	Homo	46.5	12.8	372.2	538.4	6.696
T3 3390-SP001-11-9	Homo	54.0	10.2	406.0	556.0	1026.2
T3 3390-SP001-14-2	Homo	59.7	12.5	342.4	764.0	1178.6
T3 3390-SP001-14-6	Homo	66.3	12.9	431.0	673.9	1184.1
T3 3390-SP001-15-9	Homo	30.8	14.3	271.8	559.8	876.7
T3 3390-SP001-15-12	Homo	39.6	13.1	241.7	649.1	943.5
T3 3390-SP001-16-3	Homo	49.9	17.1	230.2	519.7	816.9
T3 3390-SP001-16-6	Homo	35.5	21.1	263.8	547.7	868.1
T3 3390-SP001-35-2	Het	37.6	7.2	125.4	313.9	484.1
T3 3390-SP001-35-10	Homo	43.7	9'91	234.7	503.9	798.9
T3 3390-SP001-35-12	Homo	50.2	21.3	361.7	695.7	1128.9
T2 2200_00001_8_2	Hai	7 - 7	.0	178.7	7347	0 644
TO 2200 0000 00 CT	Llowo	30.1	10.7	300.2	0.505	7.70
1.3 3350-SF001-6-5	01110111	17.7	7.01	509.5	505.0	0/1/0
13 3390-SP001-8-11	OLIOH 	75.5	19.0	7,007	380.4	870.0
T3 3390-SP001-18-8	Het	29.5	12.2	112.1	247.6	441.1
T3 3390-SP001-16-10	Het	38.0	14.6	248.2	486.3	787.1
T4 3390-SP001-1-6-1	Homo	27.8	20.5	248.7	379.1	676.1
T4 3390-SP001-1-6-8	Homo	38.5	16.8	304.1	383.9	743.3
VAR SP001-4-5		54.2	ΩN	ΩN	5.8	0.09
VAR SP001-4-6		51.2	ΩN	ND	7.0	58.2
VAR SP001-4-10		30.2	ND	ND	Ω	30.2

Carotenoid concentration (µg/gFW)

•	Segregation	חמ					
Sample ID #	ratio		Lyconene	α-Carotene	B-Carotene	Phytoene	Total
					р-сшоши	- Iny toolic	
SP30021 control	Homo	21	ND	ND	2	ND	23
9002-SP30021-1*	3:1	20	2	394 ·	618	210	1244
9002-SP30021-2	3:1	17	2	285	537	128	969
9002-SP30021-2	>64:1	19	7	489	689	381	1585
9002-SP30021-4	3:1	58	5	105	266	94	528
9002-SP30021-5	15:1	24	3	416	649	265	1357
9002-SP30021-6	3:1	13	2	324	546	176	1061
9002-SP30021-0 9002-SP30021-7		13		344	465		_
	3:1		4	_		212	1038
9002-SP30021-8	15:1	12	3	449	690	224	1378
9002-SP30021-9	>64:1	24	5	499	724	313	1565
9002-SP30021-10	15:1	52	25	387	505	245	1214
9002-SP30021-11	3:1	29	2	301	480	187	999
9002-SP30021-12	>64:1	43	10 .	<i>5</i> 75	779	436	1843
9002-SP30021-13	3:1	19	3	357	509	279	1167
9002-SP30021-14	null	33	ND	ND	3	ND	36
9002-SP30021-15*	3:1	29	7	472	599	354	1461
9002-SP30021-16	64:1	40	3	315	436	203	997
9002-SP30021-17	15:1	25	7	322	467	144	967
9002-SP30021-18	>64:1	8	4	447	647	313	1419
9002-SP30021-19	15:1	38	17	537	570	327	1489
9002-SP30021-20*	3:1	32	8	363	629	173	1205
9002-SP30021-21	>64:1	1	6	468	736	348	1559
9002-SP30021-22	15:1	68	29	308	423	173	1001
9002-SP30021-23	15:1	51	20	449	553	423	1496
9002-SP30021-24	3:1	47	20	339	515	311	1232
9002-SP30021-25	null	27	ND .	ND	2	ND	29
9002-SP30021-26*	3:1	4	3	346	605	150	1108
9002-SP30021-27	>64:1	25	5	416	698	376	1520
9002-SP30021-28	15:1	75	9	464	527	333	1408
9002-SP30021-29	null	32	ND	16	34	ND	82
9002-SP30021-30	3:1	25	9	316	525	182	1057
9002-SP30021-31	null	28 .	ND	ND	2	ND	30
9002-SP30021-32	3:1	29	5	198	283	132	647
9002-SP30021-33	15:1	<i>5</i> 0	40	408	557	324	1379
9002-SP30021-34	15:1	43	5	216	289	132	685
9002-SP30021-35	3:1	29	8	303	511	281	1132
9002-SP30021-36	3:1	26	9	324	402	157	918
9002-SP30021-37	3:1	34	11	263	418	143	869
9002-SP30021-39	15:1	54	13	219	420	118	824
9002-SP30021-40	15:1	30	7	382	716	235	1370
9002-SP30021-41	3:1	52	15	440	506	396	1409
9002-SP30021-42	3:1	49	20	317	516	170	1072
9002-SP30021-44	>64:1	34	7	368	647	310	1366
9002-SP30021-45	>64:1	45	. 9	429	636	402	1521
9002-SP30021-45	3:1	100	14	456	699	347	1617
9002-SP30021-48		37	5	191	354	231	818
	3:1		22	522	756	303	1654
9002-SP30021-50	64:1	51	<u> </u>	J <i>LL</i>			

FIGURE 14

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	CGT	GCA	8	06	CTC	GAG	Ç		ATT	TAA	Á	180	AAC	TTG	ź		GCT	CGA	A>	270	GGT	CCA	Ĝ
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4	GTC	CAG	>		GGC	CCG		73	ACT	TGA	E		ATG	TAC	Σ	22	TTG	AAC.	ü		GGT	CCA	ပ
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	GCT	CGA	Ø	70	GTG	CAC	>		AAG	TTC	×	. 160	AGA		œ		GTT TTG	AAC	ы	250	CAC	\mathtt{GTG}	H
	TCC	AGG	Ω		GCA	CGT	Æ			CAC	>	16	GGA	CCT	Ö		GTT	CAA	>	2	ATG	TAC	Σ
20	TCT	AGA	Ω.		၁၁၅	CGG	Ø	110	CCA	GGT	<u>а</u>		GGT	CCA	ပ	200	ACC	TGG	H		ATC	TAG	Н
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	GCT	CGA	4	50	TCT	AGA	ഗ		TCC	AGG	S	140	TCC	AGG	ഗ		TTG	AAC	ᄓ	230	TCC	AGG	ഗ
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	TCT	CAA AGA	Ø	07	GAC	CTG	Ω		ACC	TGG	Ę
	GTT	CAA	>	610	CAA	GTT	ø		AGA	TCT	ជ
000	TGL	ACA	ပ		AAG	TTC	×	650	GAG	CTC	田
4.7	CAC	GTG	Ħ		TTG	AAC	ü	•	CAA	GTT	Ø
	GAC	CTG	Ω	900	AAG	TTC	×		GCT	CGA	Æ
Ç	AGA	TCT	æ		GAT	CTA	Ω	10	GAG	CTC	ជ
りなり	GGT	CCA	Ö		GTT	CAA	>	640	GCT	CGA	4
	GAG	CTC	闰	590	CCA	GGT	ы		AGA	TCT	ĸ
	GTC	CAG	>	u)	CCA	GGT	൧ഄ		$\mathbf{T}\mathbf{T}\mathbf{G}$	AAC	ы

CGT GCA XV	90 CTC GAG L>	ATT TAA I>	180 GCT CGA A>	GTC CAG V>	270 GCT CGA A>	GCT CGA
40 AGC TCG S	ပ္ပ်ပ္ပ ၅ ၅	30 GAC CTG D	TCC AGG S	220 G ATC C TAG	CAC GTG H	10 ATC TAG I
GTC CAG	ည္သ ဗ ဗ	13 ACT TGA T	ATG TAC M	TTG AAC	GTT CAA V	GCT CGA
ACA TGT	80 TTC AAG	AAC TTG N	170 TGC ACG C	TCC AGG	260 CAC GTG H	TTG AAC L
ACA TGT T	CCA GGT P	GTC CAG V	170 AAG TGC TTC ACG K C	ACC TGG	TTG	ATC TAG I
30 GTG CAC	GCT	120 AAG TTC K	GTA CAT	210 GCT CGA	GCT CGA	300 CCA GGT
GCT CGA A	70 GTG CAC V	GTG AAG CAC TTC V K	50 AGA TCT R	ACT TGA T	250 TGG TTG ACC AAC	CAC GTG
TCC AGG S	GCA CGT	GTG CAC V	160 GGA A CCT T	TTG	TGG ACC	GCT CGA
20 TCT AGA S	950 066 A	110 CCA GGT P	GGT CCA G	200 GAC CTG	GCT CGA A	290 GCT CGA
TCC AGG S	TCC AGG S	TTC	AAT TTA N	GCT CGA	GCT CGA A	GCT
ATA TAT I	60 CAA GTT Q	GGA CCT G	150 AGC TCG S	AAG TTC K	240 ATC TAG I	GAC CTG D
10 ATG TAC M	999 CCC G	100 G ACT C TGA	ACA TGT	190 G CCA C GGT	ÀTC TAG I	280 C TTG G AAC
TCT AGA S	AGG TCC R	ATG TAC M	ATT TAA I	TTG AAC L	GGT CCA G	28 TTC AAG
GCT CGA	50 TCT AGA S	TCC AGG	140 TCC AGG	GCT CGA	230 GGT CCA	TGG ACC
ATG TAC M	950 CGG	aaa TTT K	ACT TGA T	CAC GTG H	TCC AGG S	TTG
						•

	,	•																								
360	ATC	A A		AGA	TCT	2	450	GGT	CCA	&		AGA	$ ext{TCT}$	₩	540	CCA	GGT	%		AGA	TCT	\$	630	ATC	TAG	۲ ۲
	ATC	βĤ	00	CCA	GGT	ф		GCT	CGA	K	0	CAC	GTG	H		GGT	CCA	Ö	0 00	TGG	ACC	3		TTG	AAC ,	-
	TTC	F F	400	AGA	\mathbf{TCT}	æ			ATG		49		GTG			GGT	CCA	Ö	580	GGT	CCA	ტ		GCT	CGA	₹
350	TTG	1		GGT	CCA	C)	440	$\mathbf{T}\mathbf{T}\mathbf{G}$	AAC	IJ		GCT	CGA	Æ	530	CAC	GTG	X		TTC	AAG	[zı	620	TAC	ATG	H
ന	GGT	វី ច		CCA	GGT	ρı	7	TGG	ACC	3		ATG	TAC	Σ	u,	GAC	CTG	Q		TAC	ATG	×	v	GTT	CAS:	>
	GTC		390	GTC	CAG	>		TTG	AAC	ឯ	480	CAC	GTG	H		TTC	AAG	দ্র	570	ACT	TGA	٤٠		ACC	TGG E	:
10	TTG TCT	S		GTT	CAA	>	30	GTT	CAA	>		AAG	CAA TTC	×	50	GAT	CTA	Ω		GGT		Ö	10	GTT	CAA:	>
'n.	TTG	i i		TCC	AGG	ഗ്യ .	430	TTG	AAC	ы		GTT	CAA	>	520	CCA	GGT	Ωι		ATC	TAG	Н	610	ATC	TAG	-
	TGG) Z	380	GGT	CCA	ෆ		CAA	GTT	a	470	ATC	TAG					Ω	560	TTC	AAG	ĮĽι		GTC	CAG	>
	AAC	2	,	CAC	GTG	H		GGT	CCA	Ō	4	ATG	TAC	Σ		GAC	CTG	Ω	•	AGA	TCT	K		CCA	GGT	Ն լ
330	TTG	ı	•	ATG	TAC	X	420	ATG	TAC	X		AAG	$T^{T}C$	×	510	GAT	CTA	Ω		GCT	CGA	Ø	009		AAC •	
	GGT	່ວ	0,	GCT	CGA	Æ		GCT	CGA	Æ	09	AGA	$ ext{TCT}$	æ		ACT	TGA	E	550	TAC	ATG	≯		TTG	AAC	4
	TTG	ı	37	GA	CTG	D		GCT	CGA	æ	4 (TGG	ACC	Z		GGT	CCA	Ö		TGG	ACC	3		TTG	AAC	4
320	TTC	F	•	CAC	GTG	出	410	AAC	TTG	Z		TCT	AGA	ഗ	500	GCT	CGA	Ø		AGA	TCT	ĸ	590	GGT	S S S S S	פ
(°)	AAC	Z		GCT	CGA	4	7	GCT	CGA	Æ		TTC	AAG	ſĽι	4,	CAC	GTG	Ħ		GTT	CAA	>	u,	GAG	ဂ ဂ် ဂ	긔
					•												-			٠						

TCC	AGG S>	720 CCA GGT P>	CGA GCT R>	810 CAC GTG H>	CCA GGT PV	900 TAG ATC *>
CCA	GGT . P	TTG AAC L	60 GCT CGA	TTC AAG F	SO GTC CAG	GCT CGA
TTG C	AAC L	TGG ACC W	AAC TTG N	TGT ACA C	850 ACT G TGA C	ACC TGG
CCA	GGT P	710 ACC TGG	CAC GTG H	800 ACC TGG	CCA GGT	890 GAC CTG D
TGG	ACC W	GGT	AGA TCT R	TTG AAC L	CAC GTG H	GGT CCA G
TTC	AAG F	TTC AAG F	750 GAC CTG D	TTG AAC L	840 TTG AAC L	AAG TTC K
GTC	CAG V	GTT CAA V	CCA GGT P	TCC AGG	CAC GTG H	880 A ACC T TGG
GTT	CAA <	700 TTC G AAG C	TTC AAG F	790 GTT TCC CAA AGG V S	CAC GTG H	88 AGA TCT R
TAC	ATG Y	TTG AAC L	740 GCT CGA	CCA GGT P	830 GAG CTC	ACC TGG
ATG	TAC M	CAA	GAC CTG	GAT CTA D	CAC GTG	TCC AGG S
TGG	ACC	690 ATC TAG TAG	CAC GTG H	780 TCT AGA S	CAC GTG H	870 CCA GGT
C AGA	TCT R	TCT AGA S	GGT CCA G	AGA ATC TCT TAG R I	820 T TAC A ATG	TTG AAC L
GAC	CTG D	GCT CGA A	730 CCA G GGT C	AGA. TCT	GGT CCA G	AGA TCT R
GGT	င်င်နှ ရ	680 TTG AAC L	AGA TCT R	770 TCC AGG	GGT CCA G	860 TGG ACC
TTG	AAC L	ATC TAG I	CAC GTG H	TCC AGG S	TTC AAG F	TGG ACC W

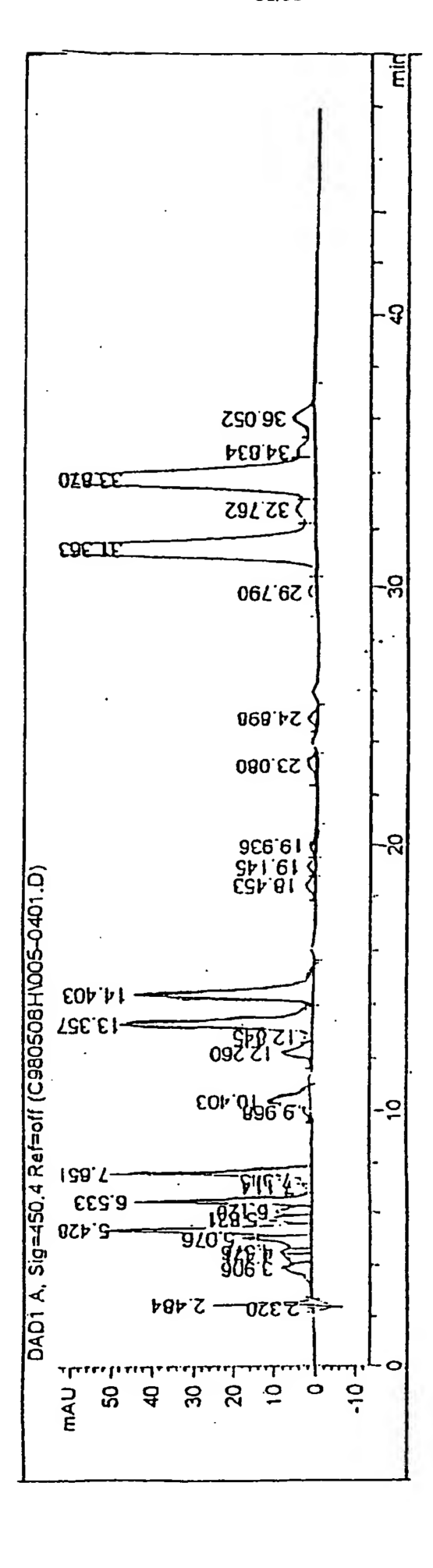
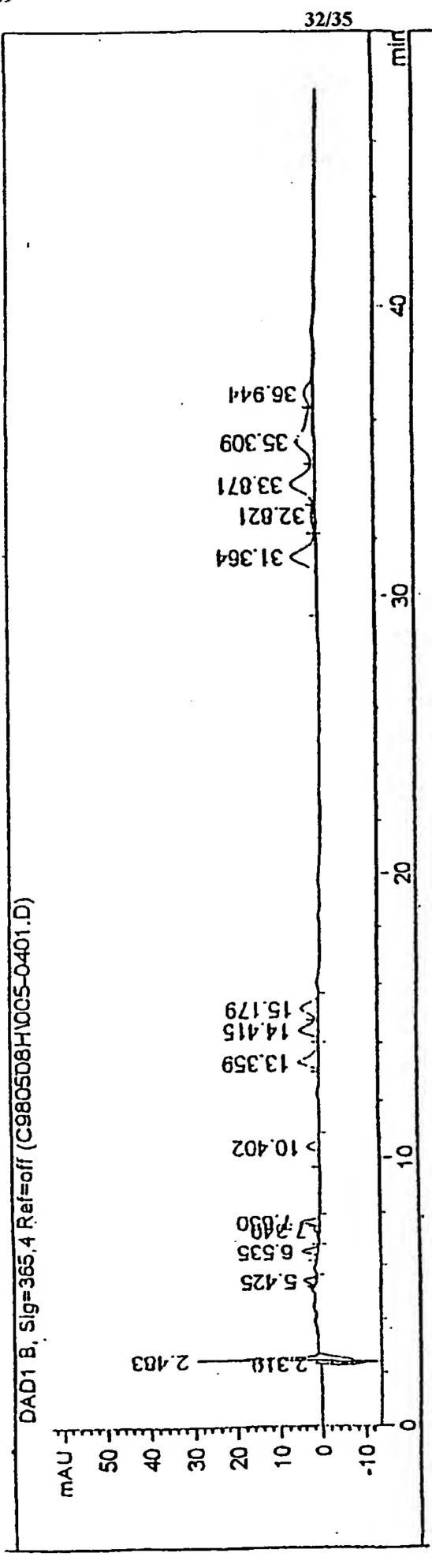
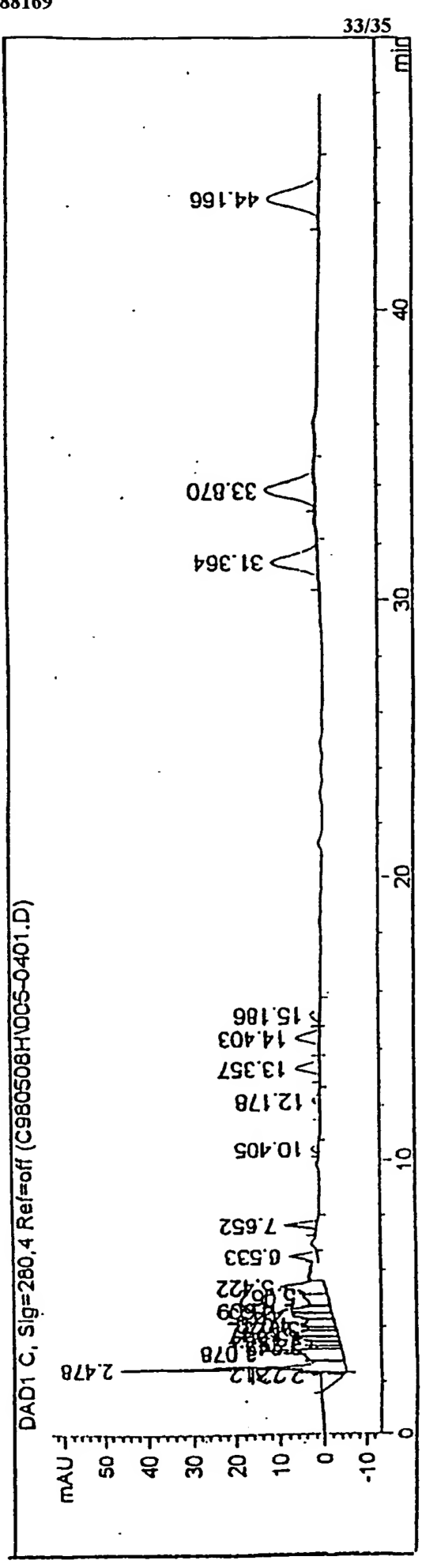


FIGURE 17A









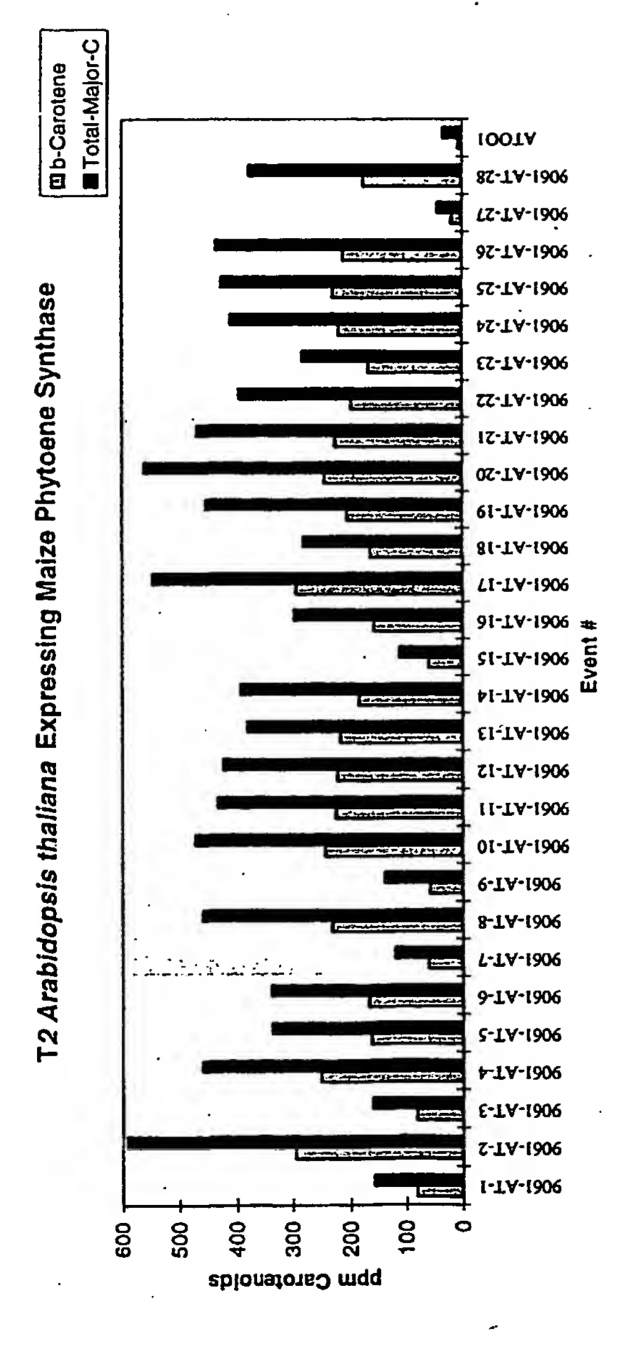


Figure 18



1	AATTCGCCCT	TCCTCCTCGA	GCGGGATCCA	TGGCCATCAT	ACTCGTACGA
51	GCAGCGTCGC	CGGGGCTCTC	CGCCGCCGAC	AGCATCAGCC	ÀCCAGGGGAC
101	TCTCCAGTGC	TCCACCCTGC	TCAAGACGAA	GAGGCCGGCG	GCGCGCCGGT
151	GGATGCCCTG	CTCGCTCCTT	GGCCTCCACC	CGTGGGAGGC	TGGCCGTCCC
201	TCCCCCGCCG	TCTACTCCAG	CCTCGCCGTC	AACCCGGCGG	GAGAGGCCGT
251	CGTCTCGTCC	GAGCAGAAGG	TCTACGACGT	CGTGCTCAAG	CAGGCCGCAT
301	TGCTCAAACG	CCAGCTGCGC	ACGCCGGTCC	TCGACGCCAG	GCCCCAGGAC
351	ATGGACATGC	CACGCAACGG	GCTCAAGGAA	GCCTACGACC	GCTGCGGCGA
401	GATCTGTGAG	GAGTATGCCA	AGACGTTTTA	CCTCGGAACT	ATGTTGATGA
451	CAGAGGAGCG	GCGCCGCGCC	ATATGGGCCA	TCTATGTGTG	GTGTAGGAGG
501	ACAGATGAGC	TTGTAGATGG	GCCAAACGCC	AACTACATTA	CACCAACAGC
551	TTTGGACCGG	TGGGAGAAGA	GACTTGAGGA	TCTGTTCACG	GGACGTCCTT
601	ACGACATGCT	TGATGCCGCT	CTCTCTGATA	CCATCTCAAG	GTTCCCCATA
651	GACATTCAGC	CATTCAGGGA	CATGATTGAA	GGGATGAGGA	GTGATCTTAG
701	GAAGACAAGG	TATAACAACT	TCGACGAGCT	CTACATGTAC	TGCTACTATG
751	TTGCTGGAAC	TGTCGGGTTA	ATGAGCGTAC	CTGTGATGGG	CATCGCAACC
801	GAGTCTAAAG	CAACAACTGA	AAGCGTATAC	AGTGCTGCCT	TGGCTCTGGG
851	AATTGCGAAC	CAACTCACGA	ACATACTCCG	GGATGTTGGA	GAGGATGCTA
901	GAAGAGGAAG	GATATATTTA	CCACAAGATG	AGCTTGCACA	GGCAGGGCTC
951	TCTGATGAGG	ACATCTTCAA	AGGGGTCGTC	ACGAACCGGT	GGAGAAACTT
1001	CATGAAGAGG	CAGATCAAGA	GGGCCAGGAT	GTTTTTTGAG	GAGGCAGAGA
1051					ATGGGCTTCC
1101					ACGACTACAA
1151		AAGAGGGCGT			
1201		ATATGGAAAA			
1251	GGCCAGACCT	AGCCACCAGA	GAAGCTGCAG	GATCCTCCTC	GAGACTGAAG
1301	GGCG				

MAIILVRAASPGLSAADSISHQGTLQCSTLLKTKRPAARRWMPCSLLGLHPWEAGRPSPAVYSSLAVNPAGEAVVSSEQK VYDVVLKQAALLKRQLRTPVLDARPQDMDMPRNGLKEAYDRCGEICEEYAKTFYLGTMLMTEERRRAIWAIYVWCRRTDE LVDGPNANYITPTALDRWEKRLEDLFTGRPYDMLDAALSDTISRFPIDIQPFRDMIEGMRSDLRKTRYNNFDELYMYCYY VAGTVGLMSVPVMGIATESKATTESVYSAALALGIANQLTNILRDVGEDARRGRIYLPQDELAQAGLSDEDIFKGVVTNR WRNFMKRQIKRARMFFEEAERGVNELSQASRWPVWASLLLYRQILDEIEANDYNNFTKRAYVGKGKKLLALPVAYGKSLL LPCSLRNGQT*PPEKLQDPPRD*RA

Figure 19